(12)

Europäisches Patentamt

European Patent Office Office européen des brevets



(11) EP 1 104 803 A1

EUROPEAN PATENT APPLICATION published in accordance with Art. 158(3) EPC

D14

- (43) Date of publication; 06.06.2001 Bulletin 2001/23
- (21) Application number: 00937223.6
- (22) Date of filing: 14.06.2000

- (51) Int Cl.7: **C11D** 11/00, C11D 11/02, C11D 17/06, C11D 3/37, C11D 3/04. C11D 3/10
- (86) International application number: PCT/JP00/03856
- (87) International publication number: WO 00/77148 (21.12.2000 Gazette 2000/51)
- (84) Designated Contracting States:
 AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
 MC NL PT SE
- (30) Priority: 14.06.1999 JP 16713999 04.04.2000 JP 2000102792 04.04.2000 JP 2000102793 02.05.2000 JP 2000133283
- (71) Applicant: Kao Corporation Tokyo 103-8210 (JP)
- (72) Inventors:
 - TAKANA, Shuji Kao Corp., Research Laboratories Wakayama-shi, Wakayama 640-8580 (JP)
 TAKAYA, Hitoshi, Kao Corp., Research Laboratories
 - Wakayama-shi, Wakayama 640-8580 (JP)

- YAMABOSHI, Hiroki Kao Corp., Research Laboratories
 Wakayama-shi Wakayama 640-8580 (JP)
- SUGIYAMA, Yoichi Kao Corp.,
 Research Laboratories
- Wakayama-shi, Wakayama 640-8580 (JP)

 KITAGAITO, Hiroshi Kao Corp.,
- Research Laborator.
- Wakayama-shi Wakayama 640-8580 (JP)
- YAMAGUCHI, Shu Kao Corp., Research Laboratories Wakayama-shi Wakayama 640-8580 (JP)
- YAMASHITA, Hiroyuki Kao Corp., Research Labor.
 Wakayama-shi Wakayama 640-8580 (JP)
- (74) Representative: HOFFMANN EITLE Patent- und Rechtsanwälte Arabellastrasse 4

81925 München (DE)

- (54) GRANULES FOR CARRYING SURFACTANT AND METHOD FOR PRODUCING THE SAME
- (57) The present invention relates to particles for supporting a surfactant and a process for preparing the same. Further, the present invention relates to high-density detergent particles using the particles for supporting a surfactant, and a process for preparing the same. According to the present invention, there can be obtained particles for supporting a surfactant which are excellent in the supporting ability (supporting capacity/supporting strength) for the liquid surfactant composition, and particles for supporting a surfactant which are excellent in the absorption properties (supporting rate) for the liquid surfactant composition, by spray-drying a preparation liquid obtainable by a process comprising the steps of preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble polymer and a water-soluble salt; and subjecting the first preparation liquid to a treatment of increasing a number of water-soluble salt particles, thereby preparing a second preparation liquid having an increased number of water-soluble

salt particles, as compared to the number of water-soluble salt particles which are present in the first preparation liquid. Further, since the liquid surfactant composition is supported by the particles for supporting a surfactant, detergent particles which are excellent in detergent performance, quality and the like can be efficiently obtained.



FIG. 4

Description

TECHNICAL FIELD

[0001] The present invention relates to particles for supporting a surfactant, and a process for preparing the same. Further, the present invention relates to high-density detergent particles using the particles for supporting a surfactant, and a process for preparing the same.

BACKGROUND ART

10

[0002] One process for obtaining a powdery delergent includes a process comprising the step of supporting a liquid surfactant. In this process, a high supporting alliquid surfactant is demanded for the particles for supporting a surfactant. In other words, there are two factors for the supporting a surfactant. A large amount of a liquid surfactant can be retained (supporting demanded for the particles for supporting a surfactant. A large amount of a liquid surfactant can be retained (supporting demanded for the particles for supporting surfactant can be supporting retained in the inner portion of the particle without being bleeded out (supporting strength). The supporting capacity is important from the viewpoint of formulating a surfactant in an amount necessary for detergency performance, and the supporting strength is also important from the viewpoints of preventing lowering the flowability of powdery detergent, caking, and migration of the liquid surfactant to a container or its surface by supports gondernous and container or its support g

[0003] Further, from the viewpoint of productivity, a property of quickly absorbing the liquid surfactant (supporting rate) is also demanded for the particles for supporting a surfactant.

[0004] As to the structure demanded for the particles for supporting a surfactant having a high supporting ability, it is desired to have a structure so that the supporting eacachy is increased by having a sufficient microprorus capacity in the inner portion of the particle, and that the supporting strength is high by having fine micropror diameter. Such a structure is obtained by constructing the particles for supporting a surfactant with the particles are in contact with each other, with maintaining a sufficient air gap therebetween. As a supplying source for the integrations, and the supplying source for the integrations are sufficient to sufficient solutions. It is not sufficient to sufficient sufficient solutions are sufficient to sufficient su

[9005] As a technique for actualizing such formation, Japanese Patent Laid-Open No. Sho 62-112697 discloses a process of obtaining a dry powder having a high absorption capacity (particles for supporting a surfactant), comprising adding and mixing a crystal growth-controlling agent, which is an organic substance having at least 3 carboxyl groups in the molecule, in an effective amount, with a sturry, prior to mixing the sturry with sedium carbonate, thereby forming socilum carbonate monohydrate and/or burkeite, of which crystal growth is controlled, in the sturry, and thereafter supprise drying the mixture sturry.

[0006] However, the supporting ability of the particles for supporting a surfactant obtained by this process has not been sufficient. The causes therefor include the amount of the fine burkeite dispersed being insufficient in the sturry before soray-drying; and the amount of the fine scicular crystals of burkeite being insufficient also in the particle obtained by spray-drying. The fine burkeite crystals are a base material effective for improving the supporting ability. However, in this technique, since dissolved sodium suffate forms burkeite on the surface or neith surface of granular sodium carbonate added afterwards, a majority exists as an aggregate which is hard and has a large particle size. Therefore, the amount of the burkeite in a fine accidat crystal state formed in the sturry is small, and the burkeite which could have been inherently formed into fine acciduar crystals takes an aggregated state having a large particle exise in the particle exist. Therefore, the resulting particles have lavel great great shay large microprocus capacity and micropore.

[0007] Also, a polyacrylate (polymer), which is a polymer especially effective as a crystal growth-controlling agent, may form a coating film on the particle surface. Therefore, when the polymer is formulated as a detergent composition in an effective amount or more, there may be some cases where the resulting particle does not exhibit a sufficient supporting ability. In this publication the maximum supporting capacity is exhibited when the amount of the polymer in the particle is as small as about 1 to about 2% by weight, so that a certain limitation must have been added to the formulation amount of the water-soluble polymer.

diameter, so that a sufficient supporting ability cannot be exhibited.

[0008] The water-soluble polymer is a base material having a film-forming characteristic by drying. When the watersoluble polymer is formulated in the slurry, a coating film containing a water-soluble polymer on a particle surface after drying is formed, thereby lowering the degree of porocity. In this case, the supporting rate tends to be lowered, so that a certain period of time has been required for sufficiently supporting a fliquid surfactant in the particles for supporting a surfactant. In order to efficiently prepare detergent particles by the process of supporting the fliquid surfactant in the

particles for supporting a surfactant, it has been desired to further increase the supporting rate for the liquid surfactant composition in the particles for supporting a surfactant.

DISCLOSURE OF INVENTION

5

15

20

25

30

35

[0009] Accordingly, an object of the present invention is to provide particles for supporting a surfactant which are excellent in the supporting ability (supporting capacity/supporting stergingly) of the liquid surfactant composition; a process for preparing the particles for supporting a surfactant, particles for supporting a surfactant which are excellent in the absorption property (supporting rate) of the liquid surfactant composition; detergent particles prepared by using the particles for supporting a surfactant is detergent composition comprising the detergent particles; and a process for preparing detergent particles prepared by using the particles for supporting a surfactant.

[0010] These objects and other objects of the present invention will be apparent from the following description.

[0011] Specifically, the present invention relates to:

[1] a process for preparing particles for supporting a surfactant comprising the steps of preparing a preparation licuid comprising a water-soulble polymer and a water-soluble sell, and spray-drying the preparation liquid obtained thereby, wherein the step of preparing the preparation liquid comprises (a) preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble polymer and a water-soluble salt, and (b) subjecting the first preparation liquid to a treatment of increasing a number of water-soluble salt particles, thereby preparing a second preparation liquid to alwing an increased number of water-soluble salt particles, as compared to the number of water-soluble salt particles which are present in the first preparation liquid.

[2] particles for supporting a suffactant obtainable by spray-drying a preparation liquid comprising a water-soluble polymer and a water-soluble sail, wherein the particles for supporting a surfactant have a mode diameter of the microprovus capacity distribution, as determined by mercury pronsimeter, of 1.5 µm or less, a microprovus capacity of 0.3 mU/g or more for one having a micropore diameter of from 0.01 to 3.0 µm, and a particle strength of from 15 to 100 Mb/a:

[3] particles for supporting a surfactant comprising a water-soluble polymer and a water-soluble salt, wherein at least a part of particles comprises a particle which is a cave-in particle having a structure that there exists a hollow, namely a cave-in hole, in an inner portion thereo, and that a particle surface is opened and communicated the

the hollow in the inner portion;

[4] a process for preparing detergent particles having a bulk density of from 500 to 1000 g/L, comprising the step of mixing from 10 to 100 parts by weight of a surfactant composition with 100 parts by weight of particles for supporting a surfactant obtainable by the process of Item [1] above or the particles of Item [2] above,

[5] detergent particles having a bulk density of from 500 to 1000 g/L, wherein from 10 to 100 parts by weight of a strictant composition is supported in 100 parts by weight of particles for supporting a surfactant obtainable by the process of item [1] above or the particles of item [2] above; and

[6] a detergent composition comprising the detergent particles of item [5] above.

BRIEF DESCRIPTION OF THE DRAWINGS

[0012] Figure 1 is a show of an SEM photograph showing one example of an external appearance of the particles for supporting a surfactant comprising a cave-in particle.

[0013] Figure 2 is a show of an SEM photograph for a split cross section of the cave-in particle.

[0014] Figure 3 is a schematic view of the particle observed from the surface centering about a cave-in hole.

[0015] Figure 4 is a schematic side view of a cross section obtained by perpendicularly splitting the particle against the face centering about a cave-in hole as shown by a broken line in Figure 3.

BEST MODE FOR CARRYING OUT THE INVENTION

50 1. Definitions of Terminology

[0016] The term "particle for supporting a surfactan" of the present invention refers to a particle obtainable by spraydrying a preparation liquid comprising a water-soluble polymer and a water-soluble salt, which is used for supporting a liquid surfactant composition, and an aggregate thereof is referred to as "particles for supporting a surfactant." The term "detergent particle" refers to a particle comprising a surfactant, a builder and the link, in which a liquid surfactant composition is supported in a particle for supporting a surfactant, and the term "detergent particles," means an aggregate thereof. The term "detergent composition" means a composition comprising detergent particles, and further comprising separately added detergent components other than the detergent particles as desired (for instance, builder particles,

fluorescent dyes, enzymes, perfumes, defoaming agents, bleaching agents, bleaching activators, and the like). In the present specification, the preparation liquid may be referred to as a "first preparation liquid" and a "second preparation liquid" in some cases. The second preparation liquid is obtained by treafing the first preparation liquid control of the preparation liquid in some cases. The second preparation liquid in some undissolved substances and precipitates derived from a water-solube salt. The term "undissolved substances" means a water-solube salt, which cannot be dissolved in the liquid phase and is present as a solid, among raw materials added to the first preparation liquid, and the term "precipitate" means a solid derived from a water-soluble salt fromed from a liquid phase of the first preparation liquid. Also, in the phrase "derived from a water-soluble salt fromed from a liquid phase of the first preparation liquid. Also, in the phrase "derived from a water-soluble salt fromed from a liquid phase of the first preparation liquid. Also, in the phrase "derived from a water-soluble salt of the first preparation liquid. So, in the phrase "derived from a water-soluble salt of the first preparation liquid. So, in the phrase "derived from a water-soluble salt of means a water-soluble salt of the solublity is of the phrase "derived from a water-soluble salt of means a water-soluble salt of the solublity is solublity is set than 0.5 of 100 g of water at 25°C, and molecular weight is 1000 or more. The term "liquid surfactant composition" refers to a composition comprising a liquid or paste-like surfactant when supporting the surfactant in the particles for supporting a surfactant.

2. Improvement in Supporting Ability of Particles for Supporting Surfactant

15

[0017] The properties required for the particle for supporting a surfactant (hereinather also referred to as "particle for supports") to estibilit a high supporting ability include having much space (supporting sites) for supporting a liquid surfactant composition (hereinather also referred to as "liquid composition") in the inner portion of the particle, namely having a large microporous capacity for the inner portion of the particle, thereby having a large supporting capacity for the liquid composition, and having a small micropore diameter in the inner portion of the particle, thereby having strong supporting strength for the liquid composition. In addition, it is necessary that the particle for supports has a high supporting rate for the liquid composition for effectively making use of the supporting sites in the inner portion of the particle, and has a particle strength durable when preparing a detergent particle for the operation such as mixing for supporting the fluid composition.

[0018] In the particles for supporting a surfactant obtainable by spray-drying a preparation liquid comprising a watersoluble polymer and a water-soluble salt, a method for improving a supporting rate with dramatic improvements in the supporting ability and the particle strength has been studied. As a result, completely new facts not conventionally touch have been found that the microporous capacity of the inner portion of the particle obtainable by spray-drying the preparation liquid can be made even larger, and the micropore diameter of the inner portion of the particle can be made smaller by increasing the number of the water-soluble salt particles which are present in the preparation liquid to be spray-dried, and that the formation of the coating lift in the particle surface is suppressed.

[0019] As the number of the water-soluble salt particles which are present in the preparation liquid increases, the particles are present as dispersion of fine particles in the preparation liquid. In addition, the fine particles are present in a dispersed state in the inner portion of droplets in the process of spray-drying the droplets of the preparation liquid. As described above, the fine water-soluble salt particles which are present in a dispersed state in the inner portion of the spraying droplets contribute to the formation of the supporting sites by being retained in a dispersed state even in the inner portion of the particle obtainable by spray-drying. In other words, the water-soluble salt which is present in the preparation liquid has a large specific surface area by increasing its number, and is utilized in a more effective formation of the supporting site for the liquid composition in the particle obtainable by spray-drying. Further, it has been found that there may be some cases where the fine water-soluble salt particles play a role as seed crystals when the water-soluble salt dissolved in the liquid phase of the preparation liquid is precipitated in the spray-drying process. Here, the fine water-soluble salt particles can serve as seed crystals, when the fine water-soluble salt particles contain the same salt as the water-soluble salt and/or a compound salt of the salt which is dissolved in the preparation liquid and/or a solid of a complex salt. Moreover, in the spray-drying process, the water-soluble salt dissolved in the liquid phase of the preparation liquid is precipitated as fine acicular crystals subjected to crystal growth adjustment action of the water-soluble polymer by having the seed crystals dispersed in the inner portion of the spray droplets as a core, thereby more effectively making use of improvements of the supporting sites of the inner portion of the particle. Since the particles for supports obtainable by this process can have small micropore diameter of the inner portion of the particle, they are excellent in the supporting ability, especially supporting strength, for the liquid composition, and have high particle strength.

[0020] The technique for improving the supporting ability of the particles for supporting a surfactant described above is effective when preparing both a phosphorus-containing detergent containing a phosphate, and a phosphorus-free detergent, and is a technique exhibiting especially high effects when preparing a phosphorus-free detergent which is more difficult to achieve the improvement in the supporting ability.

[0021] Incidentally, the internal structure of the particle for supporting a surfactant of the present invention can be confirmed by using a mercury porosimeter as expressed by the microporous capacity distribution of the particles for

supports. In the distribution of the microporous capacity per micropore diameter of the inner portion of the particle for supports as determined by mercury porosimeter (for instance, "manufactured by SHIMADZU CORPORATION, "SHIMADZU Poresizer 9320") (hereinafter referred to as "microporous capacity distribution"), the larger the microporous capacity, the larger the supporting capacity for the liquid composition; the smaller the micropore diameter, the higher the ability of retaining a liquid composition on the aborder by capillary phenomenon (supporting strength). Therefore, in a case where the microporous capacity is even larger and a micropore diameter is even smaller, the supporting ability for the surfactant can be made high, thereby supporting a large amount of the fluid composition, and at the same time the bleed-out of the liquid composition can be suppressed. Therefore, the particles for supporting a surfactant of the present invention which are suitable for supporting the fluid documposition to a mode diameter of the microporous capacity distribution (the micropore diameter having the largest microporous capacity in the obtained microporous capacity distribution) of 1.5 µm or less, preferably 1.3 µm or less, more preferably 1.1 µm or less, still more preferably 1.0 µm or less, specially preferably 0.9 µm or less, most preferably 0.8 µm or less, so

10022] In addition, with regard to the microporous capacity of the particles for supporting a surfactant of the present invention, the microporous capacity of one having a micropore diameter of from 0.01 to 3.0 µm is 0.3 mHz or more. It is preferable that the microporous capacity of one having a micropore diameter of from 0.01 to 2.5 µm is 0.3 mHz or more. It is smore preferable that the microporous capacity of one having a micropore diameter of from 0.01 to 2.0 µm is 0.3 mHz or more. It is still more preferable that the microporous capacity of one having a micropore diameter of from 0.01 to 1.5 µm is 0.3 mHz or more. It is still more preferable that the microporous capacity of one having a micropore diameter of from 0.01 to 1.0 µm is 0.3 mHz or more. In addition, in the ranges of each microporous capacity is 0.3 mL/g or more, and it is still more preferable that its microporous capacity is 0.4 mHz or more.

[0023] The particle strength of the particles for supporting a surfactant of the present invention is from 5 to 200 MPa, preferably from 10 to 150 MPa, more preferably from 15 to 100 MPa, particularly preferably from 20 to 80 MPa, especially preferably from 25 to 80 MPa, from the viewpoint of preventing undesirable lowering of the supporting capacity caused by disintegration of the particle constituting the particles when the liquid surfactant composition is added to the particles. Here, the particle strength can be determined by the method described in the measurement method of the particles described below.

[0024] It is even more preferable that the particles for supporting a surfactant of the present invention have both of the above-described preferable microporous capacity distribution and particle strength. The preferable properties are such that the mode diameter of the microporous capacity distribution is 1.5 μm or less, that the microporous capacity of one having a micropore diameter of 0.01 to 3.0 μm is 0.3 ml/g or more, and that the particle strength is from 15 to 100 MPa. The more preferable properties are such that the mode diameter of the microporous capacity distribution is 1.1 μm or less, that the microporous capacity of one having a micropore diameter of 0.01 to 2.0 μm is 0.3 ml/g or more, and that the particle strength is from 20 to 80 MPa.

3. Method for Increasing Number of Water-Soluble Salt Particles Which Are Present in Preparation Liquid

[0025] There has been studied a treatment of increasing the number of water-soluble salt particles in the process of preparing a preparation liquid, which comprises (a) preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble salt, and (b) subjecting the first preparation liquid mentioned above to a treatment of increasing the number of water-soluble salt particles, thereby giving a second preparation liquid having an increased number of the particles, as compared to the number of the water-soluble salt particles which are present in the first preparation liquid. As a result, the following means (1) to (3) mentioned below have been found. [0025] Here, the preparation liquid subjected to a means for increasing the number of the water-soluble salt particles exemplified in (1) to (3) mentioned below is referred to as a second preparation liquid.

(1) precipitating a water-soluble salt dissolved in the first preparation liquid.

25

- (2) subjecting the water-soluble salt particles in the first preparation liquid to a wet pulverization.
- (3) adding to the first preparation liquid fine water-soluble salt particles which may be the same as or different from the water-soluble sailt in the first preparation liquid, under the conditions that the fine particles can be present without being substantially dissolved in the first preparation liquid.

[0027] In addition, a combination of two or more of the means (1) to (3) mentioned above is a preferable embodiment of the present invention.

[0028] Further, a process for precipitating a water-soluble salt dissolved in the first preparation liquid described in (1) has been studied. As a result, there have been found the following means.

(1-1) adding a microcrystal-precipitating agent to the first preparation liquid.

- (1-2) concentrating the first preparation liquid.
- (1-3) adjusting the temperature of the first preparation liquid so that a dissolved amount of the water-soluble salt dissolved in the first preparation liquid is lowered.
- 5 [0029] In addition, the precipitation of the water-soluble salt by a combination of two or more of the means (1-1) to (1-3) mentioned above is a preferable embodiment of the present invention.
 - [0030] Here, as a method for confirming the fact that the number of water-soluble salt particles in the second preparation liquid is increased from that of the first preparation liquid, there can be employed, for instance, the following in-line type powder droplet monitoring system (manufactured by LASENTEC, "TSUB-TEC M100"). The method for confirmation will be exemptified below.
 - [0031] One-thousand grams of the preparation liquid is weighed and placed in a 1-L stainless beaker, and stirred in a thermostat of which temperature is adjusted to the same temperature as that of the preparation liquid with rotating agitation impeliers with 3 propeller wings of 2 × 4 cm at a speed of 200 frints. An in-line type provider droplet monitoring system (manufacture by LASENTEC, TSUB-TEC MIOO?) is preparated at an angle of 45° to the liquid surface of the standard preparation liquid, and attached at a position 3 cm below the fiquid surface. By the arrangement, particles are always collided to the window surface when stirred. Lising "Control Interface for FRRM Vm. 6.4 Build 56° (manufactured by LASENTEC) as a software, a focus position is posited for each in linear side 0.02 mm from the window surface. When examined the control (measurement time) is 14.5 seconds, and the averaging (moving average) is taken with 10 measurements. The number of counts charity is particless) at the time of 5-minute measurement time is therefore.
 - [0032] The above measurements are taken for the first preparation liquid and the second preparation liquid, and the obtained number of counts is compared. Specifically, by having a larger number of counts for the second preparation liquid than the number of counts for the first preparation liquid, there can be confirmed an increase in the number of the water-soluble saft particles in the second preparation liquid as compared to that in the first preparation liquid.
- 9 [0033] Also, the increase in the number of counts can also be directly confirmed by using the above in-line type powder dropler monitoring system when preparing the second preparation liquid from the first preparation liquid. [0034] Here, the increased number of the water-soluble sait particles which are present in the first preparation liquid cannot be absolutely determined from the number of the water-soluble sait particles which are present in the first preparation liquid. For instance, the difference in the number of counts of the second preparation liquid from that of the first preparation liquid obtained by the above method may be preferably 500 particles/s or more.
 - [0035] Here, among the above means, it is preferable that the amount of the water-soluble salt undisorloved in the second preparation liquid (namely, precipitates derived from a water-soluble salt and/or fine water-soluble salt particles added to the first preparation liquid), which is increased by the treatment of increasing not only the number of the water-soluble salt particles which are present in the second preparation liquid (treatment of (1), 20) or combining two or more means of (1) to (3), is 3% by weight or more, based on the amount of the water-soluble salt dissolved in the first preparation liquid (treatment of (1), 20) or combining two or more means of (1) to (3), is 3% by weight or more, based on the amount of the water-soluble salt dissolved in the first preparation liquid before carrying out the above means. From the viewpoint of forming further effective supporting as in the inner portion of the particles after spray-drying, thereby improving the supporting ability, the amount is more preferably 5% by weight or more, still more preferably 40% by weight or more, most preferably 10% by weight or more. On the other had, from the viewpoints of securing the microporous capacity of the particles for supporting a surfactant obtained after spray-drying and the handleability of the second preparation liquid effer subjecting to the above means, the amount of the water-soluble salt undissolved in the second preparation liquid increased by the above means, perferably 25% by weight or less, more preferably 35% by weight or less, smore preferably 25% by weight or less, smore preferably 25% by weight or less, specific microporum and the water-soluble salt dissolved in the first preparation liquid.
 - [0036] The amount A (%) of the water-soluble salt undissolved in the second preparation liquid, which is increased by a means of increasing the amount of the water-soluble salt undissolved in the preparation liquid, is determined by measuring the content, the dissolution rate and the ratio of undissolved portion of the water-soluble salt in the preparation liquid before and after the treatment as determined by the subsequent method.
- 50 [0037] First, a content T (%) of the water-soluble salt of the first and second preparation liquids is determined by ion chromatography, or the like.
 - [0038] Also, the dissolution rate of the water-soluble salt is obtained as follows,
 - [033] A preparation liquid is filtered under reduced pressure, and a water concentration P (%) in the filtrate is determined by a far infrared ray heater-type moisture meter (manufactured by SHIMADZU CORPORATION) or the like. Further, the water-soluble salt concentration S (%) in the filtrate is obtained by ion chromatography or the like, byposing that the water-content of the preparation liquid is Q (%) and that content of the vater-soluble salt is of the water-soluble salt is offended by the following equation:

Dissolution Rate (%) =
$$\frac{(100 \times S \times Q)}{(P \times T)}$$
 (I)

[0040] However, when the above dissolution rate calculated exceeds 100%, the dissolution rate is considered as 100%. In addition, the ratio of undissolved portion V (%) is obtained by the following equation.

[0041] Supposing that the content of the water-soluble salt is T1 (%), the dissolution rate is U1 (%), and the ratio of undissolved portion is V1 (%) in the first preparation liquid, and that the content of the water-soluble salt is T2 (%), and the ratio of undissolved portion is V2 (%) in the second preparation liquid, the increased amount A (%) of the water-soluble salt undissolved in the above second preparation liquid is obtained by the following equation.

10

30

45

55

Increased Amount A (%) of Water-Soluble Salt Undissolved in Second

Preparation Liquid =
$$100 \times \frac{(T2 \times V2-T1 \times V1)}{T1 \times U1}$$
 (III)

20 [0442] In addition, in the preparation of first preparation it guide comprising a water-soluble polymer and a water-soluble salt, and subsequent teament of increasing the number of the water-soluble salt particles which are present in the first preparation liquid, the more finer the water-soluble salt particles which are present in the second preparation liquid are increased by the treatment, the smaller the micropore diameter of the particles for supports obtainable by spray-drying, whereby an effect of improving the supporting ability is increased. From this viewpoint, the average particle size of the water-soluble salt particles which are present in the second repearation liquid increased by the treatment is preferably 40 µm or less, more preferably 50 µm or less, still more especially preferably 30 µm or less, most preferably 10 µm or less, most preferably 10 µm or less.

[0043] The average particle size refers to an average particle size calculated from the particle size distribution resulting from subtracting the particle size distribution of the particles which are present in the first preparation liquid from the particle size distribution of the particles which are present in the second preparation liquid as determined by the following measurement method.

[10044] The particle size distribution of the particles which are present in the first or second preparation liquid can be determined by using the in-line type particle droplet monitoring system (manufactured by LASENTEC, "TSUB-TEC M100") which is used for the determination of the number of counts mentioned above. The average particle size of the water-soluble salt particles which are present in the preparation liquid described in the present specification is a measured value using "TSUB-TEC M100." The measurement is carried out in the same manner as the measurement for the number of counts described above except for determining the particle size distribution at the point of 5-minute determination. Here, the median code (particle size at which the cumulative number of particles is 50%) is defined as an average particle size. It is preferable that the water-soluble salt particles which are present in the second preparation liquid are those comprising solids composed of the same salt as the water-soluble salt dissolved in the preparation liquid and/or compound salts thereof, which can serve as seed crystals during precipitation in the process of spraydrying the water-soluble salt dissolved in the liquid phase of the preparation liquid. The water-soluble salt particles which can serve as seed crystals are those which can serve as a core during the precipitation of the water-soluble salt dissolved in the liquid phase of the preparation liquid in the process of spray-drying. And the water-soluble salt precipitating in the process of spray-drying with seed crystals as a core which are present in the dispersion state in the sprayed droplets is precipitated as fine acicular crystals which are subjected to crystal growth adjustment action of a water-soluble polymer, whereby it can be effectively utilized for improving the supporting sites in the inner portion of the particle. From the viewpoints of precipitating microcrystals in the inner portion of the particle for supports obtainable by spray-drying, thereby making the micropore diameter even smaller, and improving the supporting strength for the liquid composition and the particle strength, it is preferable that the water-soluble salt particles which can serve as seed crystals are very fine and large in number.

4. Acceleration of Absorption of Liquid Surfactant Composition Through Cave-In Hole

[0045]. As conditions for the particle for supporting a surfactant to exhibit high supporting ability, it is necessary that the particle has a large amount of space (supporting site) for supporting the liquid surfactant composition in the inner portion of the particle, Moreover, it is especially important that in the production of powdery detergent that the liquid

surfactant composition is quickly absorbed, from the viewpoint of improvement in the productivity.

[0046] As described above, when the preparation liquid generally comprising a water-soluble polymer and a watersoluble sail is spray-dried, since evaporation of moisture mainly takes place at the surfaces of the sprayed diroplets, the water-soluble components dissolved in the preparation liquid imgrate to the surface together with moisture with the progress of the sorray-drying, so that the particle obtained after spray-drying takes a spherical structure, of which surface is coated with a coating film mainly constituted by a water-soluble salt and a water-soluble polymer. The coaling film formed on the particle surface serves as a factor for delaying or inhibiting the absorption of the liquid surfactant composition into the inner confort of the particle.

[0047] Therefore, a method for increasing the supporting rate for the liquid surfactant composition in the particles for supports has been studied. As a result, it has been found that the absorption of the liquid surfactant composition is speeded up by changing the shape of the spray-dried particle (particle for supports). The spray-dried particle is obtained as an aggregate of a spherical particle obtained by influence of spherical or sprayed droplets, and it has been found that the absorption of the liquid surfactant composition is dramatically speeded up by poking a hole from the surface to the inner portion of the spray-dried particle in at least one location, for instance, poking a hole with a needle or the like. In other words, it has been found that the particles for supporting a surfactant having excellent supporting rate for the liquid surfactant composition can be obtained by changing the particle shape to have a cavein hole having a structure that there exists a hollow in the inner portion of the spray-dried particle, and a particle surface is opened and communicated with the hold with the inner portion (particle surface being caved-in).

[0043] As a method for efficiently preparing the particle for supports (cave-in particle) having the cave-in hole, a method for making caving-in the particle surface at the point of spray-dying has been studied. As a result, it has been found that the content of the exve-in particle in the sprey-dried particle can be dramatically increased by adjusting the composition to a particular range, and adjusting the water content of the preparation flouid and spray-drying conditions. [0043] The cave-in hote findle in the present invention will be described in further detail. The cave-in hote (incle) is basically present in at least one location of one particle. The action for sufficiently speeding up the absorption of the liquid surfactant composition is exhibited by this cave-in hole, and a plurality of cave-in holes may be present in one particle for causation such as interference of droples in the drying to them.

5. Explanation of Cave-In Particle

28

[0050] The phrase "particle which is cave-in particle having a structure that there exists a hollow, namely a cave-in hole, in the inner portion of the spray-dried particle, and that a particle surface is opened and communicated with the hollow in the inner portion" contained in the particles for supports of the present invention refers to a particle having an external appearance, for instance, as shown in Figure 1, and having a cross section as shown in Figure 2.

[0051] In addition, the preferable size of the cave-in hole in the cave-in particle contained in the particles for supports of the present invention will be defined. The projected area diameter of the particle can be obtained by photographing a particle using a microscope centering about the opening of the cavelin hole as shown in Figure 3, and calculating the projected area diameter from the equation (IV) by using the projected area (S1) of the particle measured from the photographed particle image.

Projected Area Diameter of Particle =
$$2 \times (S1/\pi)^{1/2}$$
 (IV)

[0052] In addition, the projected area diameter of the hole (cave-in hole) can be obtained by the equation (V) by using the projected area (S2) of the hole determined in the same manner as the projected area of the particle mentioned above with an opening as shown in Figure 2.

Projected Area Diameter of Hole =
$$2 \times (S2/\pi)^{1/2}$$
 (V)

[0053] Here, as the microscope for the above measurement, there can be used, for instance, a digital microscope "VH-6300" manufactured by KEYENCE CORPORATION and SEM such as a field emission scanning electron microscope "Model S-4000," manufactured by Hitsachi, Ltd. in the calculation of the projected area, there can be used, for instance, WinRoof manufactured by Mitsutani, and the like.

[0054] A preferable diameter for the hole which is present in the cave-in particle contained in the particles for supports of the present invention is a hole in which:

(Projected Area Diameter of Hole) (Projected Area Diameter of Particle) × 100

is 2% or more. In addition, from the viewpoints that the fluid surfactant composition is easily infiltrated by and ontered hrough the cave-in-hole, and that a particle shape even closer to a spherical shape is desired for external appearance, the above rallo is preferably from 2 to 70%, more preferably from 4 to 60%, still more preferably from 5 to 50%, especially preferably from 8 to 40%, most preferably from 10 to 30%.

[0055] The depth of lihe hole which is present in the cave-in particle contained in the particles for supports of the present invention is expressed by the ratio of a distance of between a tangent line X of an open surface of the cave-in hole and a tangent line Y with the bottom of the hole in parallel to the tangent line X as shown in Figure 4 to the projected area diameter of the particle described above, i.e.,

Here, the depth of the hole can be determined, for instance, by splitting a particle with a surgical knife or the like at a plane perpendicular to the open hole portion of the cave-in hole as shown by the broken line in Figure 3, and photographing the cross section with SEM or the like. It is preferable that the depth of the hole which is present in the cave-in particle contained in the particles for supporting a surfactant of the present invention is such that the ratio as defined above is 10% or more. In addition, from the viewpoints of even more increasing the supporting rate for the liquid surfactant composition and even more securing the supporting capacity for the liquid surfactant composition in the inner portion of the particle in a large amount, the ratio is more preferably from 10 to 90%, more preferably from 15 to 80%, especially preferably from 20 to 70%.

[0056] It is desired that the content of the cave-in particle in the constituent particle of the particles for supports of the present invention is 30% or more, preferably 50% or more, more preferably 70% or more, still more preferably 80% or more more and 100% or less, from the viewpoint of more speedily and effectively absorbing the liquid surfactant composition, thereby increasing the productivity.

[0057] In addition, the constituent particle other than the cave-in particle mentioned above in the present invention includes particles having a size outside that defined as the cave-in hole mentioned above, a spiit particle and a spherical particle having no cave-in holes, and the like. It is desired that the content of these constituent particles is 70% or less, preferably 50% or less, more preferably 30% or less, \$\text{sill}\$ more preferably 20% or less, most preferably 10% or less.

[0058] Here, the content of the cave-in particle in the present invention is determined by the following method. Specifically, using inne-step sleves each having a sleve-opening as defined by J.15 2 8801 of 2000 µm. 400 µm. 100 µm

6. Composition of Particles for Supporting Surfactant

15

20

- 35

45

[0059] The particles for supports of the present invention are mainly composed of a water-soluble polymer and a water-soluble sail. The water-soluble polymer and the water-soluble sail are important for forming a supporting is and a cave-in hole for a liquid surfactant composition. In addition, the water-soluble polymer has an action of imparting strength to the carticle.

[0060] The preferable water-soluble polymer can be exemptified, for instance, by one or more kinds selected from the group consisting of carboxylic acid-based polymens; cellulose derivatives such as carboxymethy celluloses; aminocarboxylic acid-based polymers such as polyglyoxylates and polyaspartates; water-soluble starches; sugars; and the like. Among them, the carboxylic acid-based polymers are preferable, from the viewpoints of the action of making the water-soluble salt fine and the detergency, concretely including the action of capturing metal ions, the action of dispersing solid particle stains from garments into a washlub, and the action of preventing the particle stains from redepositing to the garments.

[0061] Among the carboxylic acid-based polymers, acrylic acid homopolymers and the salts thereof (Na. K. NHa.

and the like), and acrylic acid-maleic acid copolymers and the salts thereof (Na, K, NH₄, and the like) are especially excellent

[0062] The weight-average molecular weight of these water-soluble polymers is preferably from 1000 to 300000, more preferably from 2000 to 100000, still more preferably from 2000 to 80000, particularly preferably from 5000 to 50000, especially preferably from 6000 to 20000.

[0063] The molecular weight is determined as follows

- 1. Standard substance for calculation: polyacrylic acid (AMERICAN STANDARDS CORP)
- 2. Eluent: 0.2 mol/L phosphate buffer/CH2CN: 9/1 (volume ratio)
- 3. Column: PWXL + G4000PWXL + G2500PWXL (manufactured by Tosoh Corporation)
- 4. Detector: RI

10

- 5. Sample concentration: 5 mg/mL
- 6. Injected amount: 0.1 mL
- 7. Temperature for determination: 40°C
- 8. Flow rate: 1.0 mL/min

[0064] In addition to the above carboxylic acid-based polymers, polymers such as polyglyoxylates; cellulose derivatives such as carboxymethyl cellulose; and aminocarboxylic acid-based polymers such as polyaspartates can be used as ones having a metal ion capturing ability, a dispersibility and an ability of preventing re-deposition.

[0065] Other polymers include polywiny pyrrolidones (FVP), polyethylene glycols (PEG), polypropylene glycols (PEG), polypropylene glycols (PEG), and the PPG having a molecular weight of from about 1000 to about 20000 are preferable, a because the viscous characteristic of a paste, which is caused by containing water of a powder deterent, is improved.

[0068] The content of the water-soluble polymer in the particles for supports is preferably from 2 to 30% by weight, more preferably from 6 to 30% by weight, still more preferably from 6 to 26% by weight, still more preferably from 8 to 26% by weight, still more preferably from 8 to 26% by weight, was preferably from 8 to 26% by weight, was preferably from 8 to 27% by weight. Within the above range, the particle has a sufficiently high strength.

[0067] The water-soluble salt includes water-soluble inorganic salts having a carbonate group, a sulfate group, a hydrogeneuflate group, a hosphate group, and the like (for instance, alkali metal salts, armonium salts, or amire salts), in addition, there may be included halides such as chionides, bromides, iodides, and fluorides of alkali metal salts (for instance, sodium or potassium salt) and alkaline earth metal salts (for instance, calcium or magnesium salt), in addition, there can be included compound salts containing these salts (for instance, burkeite, sodium esequicarbonate, and the like).

[0068] Among them, carbonates, suifates and suffates are preferable. Carbonates are preferable as an alkalizing is agent for showing a suitable pit buffering region in washing liquid, and salts having a high degree of dissociation such as suifates and sulfites enhance an ionic strength of a washing liquid, and favorably act to sebum stains. In addition, suffites reduce hypochlorite ions contained in tap water, and have an effect of preventing detergent components such as enzymes and perfumes from oxidation degradation by the hypochlorite ions.
[0069] Sodium tripolyphosphates can also be used as the water-souble salt.

[0070] The water-soluble salt may be composed of a single component, or may be a combination of a plurality of components such as a carbonate and a sulfate.

[0071] In addition, since the water-soluble sait changes its crystal structure when precipitated in the presence of a water-soluble polymer, the water-soluble sait plays an important role in the improvement of the supporting ability of the particles for supports. Among them, as a base makerial for forming the supporting sites of the particles for supports, carbonates and/or sulfates are more preferable, and especially a combination of solution carbonate and sodium sulfate is most preferable. Especially, sodium carbonate and/or burketle, which is a compound sait of sodium carbonate and addium sulfate, is important as a base material for forming the supporting sites of the particles for supports.

[0072] In addition, since halides of alkali metals and/or alkaline earth metals, such as sodium chioride, effectively form the supporting sites of the particles for supports as microcyatel precipitating agents, because they have an enfect, when added to a first preparation liquid comprising sodium carbonate and/or sodium sulfate, or dissolving themselves and in turn precipitating microcystatis of sodium carbonate or sodium sulfate, or a compound salt thereof. Further, these halides also are especially favorable because they also have an action of partially suppressing the formation of a surface coating film in the drying process, whereby exhibiting an action of increasing supporting rate for the liquid composition in the particles for supports.

15 [0073] In addition, from the viewpoints of satisfying both the supporting ability of the particles for supporting a surfactant and the deterging performance when used as a detergent composition, a preferable weight ratio of (sodium carbonate) to (sodium sulfate) in the particles for supports is from 1:0 to 1:5, more preferably from 1:0 to 1:4, still more preferably from 1:0 to 1:3, especially preferably from 1:0 to 1:1.

[0074] In addition, from the viewpoints of salisfying both the particle strength of the particles for supporting a surfactant and the deterging performance when used as a detergent composition, a preferable weight ratio of (sodium carbonate and/or sodium sulfate) to (water-soluble polymer) in the particles for supports is from 19:1 to 1:1, more preferably from 15:1 to 1:5.1, at 11 to 2:5.1, most preferably from 8:1 to 2.5:1.

[0075] În addition, a water-soluble organic salt having a low molecular weight can also be used as the water-soluble salt, and includes, for instance, carboxylates such as citrates and furnarates. In addition, from the viewpoint of the detergency, preferable ones include methylminodiacetates, iminodisuccinates, ethylendeaimnodiacucinates, furnine diacetates, hydroxylthylminodiacetates, β-alanine diacetate, hydroxylminodisuccinates, methylglycine diacetate, quitamic acid diacetate, angeragine diacetate, senter diacetate, and the like.

[0076] The content of the water-soluble salt in the particles for supports is preferably from 20 to 90% by weight, more preferably from 30 to 80% by weight, most preferably from 40 to 70% by weight. Within these ranges, the particles for supports have a sufficiently high particle strength, and the ranges are preferable from the viewpoint of the dissolubility of the deteroent particles.

[0077] In addition, the particles for supporting a surfactant of the present invention can comprise a water-insoluble substance. As the water-insoluble substance, there can be used crystalline aluminosilicates, amorphous aluminosilicates, silicon dioxides, hydrated silicate compounds, clay compounds such as pertite and bentionite, and the like. From the viewpoints of its contribution to support for the liquid surfactant composition and not promoting generation of undissolved remnants, and the like, the crystalline aluminosilicates and the emorphous aluminosilicates are preferable. In addition, the average particle size of the aluminosilicates is preferably from 0.1 to 10 µm, more preferably from 0.5.

(loo78) Preferable crystalline aluminosilicates include A-type zeoilites (for instance, trade name: "TOYOBUILDER," manufactured by Tosoh Corporation; trade name: "Gosei Zeoilite," manufactured by Mopon Builder K.K.; trade name: "WALFOR 100," manufactured by PC OFEMICALS (Thalland), Lid.; trade name: "ZEOBUILDER," manufactured by ZEOBUILDER Lid.; trade name: "Necole Note of the Total Control of the Note of the of the Note

[0079] In addition, amorphous aluminosilicates, amorphous silicas, and the like, which have a high oil-absorbing ability but a low metal ion capturing ability, can be used as the water-insoluble substances. Examples include amorphous aluminosilicates including those described in Japanese Patent Lald-Open No. Sho 62-191417, page 2, lower right column. Ilne 19 to page 5, upper left column, line 17 (especially, the initial temperature being preferably within the range from 15° to 60°C); and those described in Japanese Patent Laid-Open No. Sho 62-191419, page 2, lower right column, line 20 to page 5, lower left column, line 11 (especially, the oil-absorbing amount being 170 mL/100 g); amorphous aluminosilicates (oil-absorbing ability: 285 mL/100 g) described in Japanese Patent Laid-Open No. Hei 9-132794, column 17, line 46 to column 18, line 38; Japanese Patent Laid-Open No, Hei 7-10526, column 3, line 3 to column 5, line 9; Japanese Patent Laid-Open No. Hei 6-227811, column 2, line 15 to column 5, line 2; Japanese Patent Laid-Open No. Hei 8-119622, column 2, line 18 to column 3, line 47, and the like. For instance, there can be used oilabsorbing carriers, for instance, "TOKSIL NR" (manufactured by Tokuyama Soda Co., Ltd.; oil-absorbing ability: 210 to 270 mL/100 g); "FLOWRITE" (the same as above; oil-absorbing ability: 400 to 600 mL/100 g); "TIXOLEX 25" (manufactured by Kofran Chemical; oil-absorbing ability: 220 to 270 mL/100 g); "SILOPURE" (manufactured by Fuji Devison Co., Ltd.; oil-absorbing ability: 240 to 280 mL/100 g), and the like. Especially, as the oil-absorbing carriers, favorable are those described in Japanese Patent Laid-Open No. Hei 6-179899, column 12, line 12 to column 13, line 1, and column 17, line 34 to column 19, line 17,

[0080] The water-insoluble substance may be composed of a single component, or a plurality of components.

[0081] The content of the water-insoluble substance in the particles for supports, when the water-insoluble substance is contained therein, is preferably from 8 to 49% by weight, more preferably from 16 to 45% by weight, most preferably from 24 to 40% by weight. Within this range, the particles for supporting a surfactant excellent in the particle strength and the dissolubility can be obtained.

[0082] Especially, in the particles for supports of the present invention, it is preferable that the content of the watersoluble polymer is from 2 to 30% by weight, that the content of the water-soluble sall is from 20 to 90% by weight, and that the content of the water-insoluble substance is from 8 to 49% by weight.

[0083] As other components, a surfactant can be formulated in the particles for supports. However, in a case where the second preparation liquid comprises a surfactant, a coating film tends to be formed on the surface of the resulting particle for supports in the process of spray-drying for preparing the particles for supports. Therefore, as a result, not

only the absorption rate of the liquid surfactant composition to the particles for supports is lowered, but also the formation of the cave-in hole is indexer. Therefore, from these viewpoints, the lower the content of the surfactant in the particles for supports the better, and it is preferable that the surfactant is rather not present. From the above reasons, the content of the surfactant in the particles for supports is preferably from 0 to 3% by weight, more preferably from 0 to 1% by weight, and especially most preferably substantially not contained.

[0084] As examples of the surfactant, the same ones as those for the liquid surfactant composition to be supported in the particles for supports described below can be used.

[0885] The amorphous silicates have an action of enhancing the particle strength of the particles for supports, In a case where the particles for supports comprise a water-insoluble substance such as an aluminosilicate, when the amorphous silicate is contained in the second preparation liquid for preparing the particles for supports, aggregated lumpy masses are formed, which become slightly water-soluble with the passage of time. Therefore, it is preferable that the crystalline silicate is substantially not contained. In addition, since the crystalline silicate is sold dissolves in the second preparation liquid to become amorphous, it is also preferable in the same manner as the amorphous silicate that the crystalline silicate is not contained in the second preparation liquid. Also, in a case where a water-insoluble substance such as an aluminosilicate is not used, when the silicate is formulated in the second preparation liquid, there is exhibited a tendency of a lowered dissolution rate of the particles for supports obtained after spray-drying. Therefore, it is preferable that the amount of the silicate contained in the second preparation liquid is 10% by weight or less, more preferably 5% by weight or less, more preferably 5% by weight or less, more preferably substantially not contained, based on the water-soluble sail excluding the silicate contained in the second preparation liquid.

[0066] In addition, the particles for supports can contain auxiliary components such as fluorescent dyes, pigments, dyes and enzymes. The content of the auxiliary components in the particles for supports is preferably 10% by weight or less, more preferably 5% by weight or less.

7. Process for Preparing Particles for Supporting a Surfactant

25

30

35

55

[0087] The particles for supporting a surfactant of the present invention can be prepared by spray-drying a second preparation liquid obtained by a process comprising step (a) and step (b) described below.

Step (a): preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble polymer and a water-soluble salt; and

Step (b): subjecting the first preparation liquid to a treatment of increasing a number of water-solube salt particles, thereby preparing a second preparation liquid having an increased number of particles of water-soluble salt particles, as compared to the number of water-soluble salt particles which are present in the first preparation liquid.

1088] Here, as to the step of dryling a preparation liquid prepared by the process comprising step (a) and step (b), the second preparation liquid may be directly subjected to dryling, or as occasion channels, for instance, it may be subjected to dryling after such a process as dilution or defoaming in order to improve the handleability of the preparation liquid. As to the dryling process, all sorts of dryling processes, for instance, freeze-dryling, dryling under reduced pressure, and the like, can be employed. From the viewpoint of effectively acting the water-coluble sait particles contained in the second preparation liquid in which the number of particles is increased for supporting the liquid composition, it is preferable that the preparation liquid to be subjected to dryling is instant-dried. Therefore, an especially preferable dryling process is a spray-drying process. As to the spray-dryling lower, those of the forms of both the countencurrent tower and occurrent tower and be used, and the countencurrent tower in the viewpoint of productivity, in addition, as a heat source for the spray-dryling lower, a pulse-impulse wave dryer using a pulse combustor may be exemplified as one of preferable dryling apparatus. In the pulse-impulse wave dryer using a pulse combustor may be exemplified as one of preferable dryling apparatus. In the pulse-impulse wave dryer includes PULCON (manufactured by Osaka Fujik Kopyo Kabusnik Kaisha).

100891 Preferred embodiments of step (b), as described above, are roughly classified into:

- (1) an embodiment of precipitating a water-soluble salt dissolved in the first preparation liquid;
- (2) an embodiment of subjecting water-soluble salt particles in the first preparation liquid to wet pulverization; and (3) an embodiment of adding fine water-soluble salt particles to the first preparation liquid under conditions that
- the fine particles are capable of being present without substantially being dissolved in the first preparation liquid.

[0090] These embodiments will be described in detail below.

7-1. Precipitation of Water-Soluble Salt Dissolved in First Prenaration Liquid

[0091] This embodiment comprises (a) preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble polymer and a water-soluble salt; and (b) precipitating a water-soluble salt dissolved in the first preparation liquid. The water-soluble salt precipitated in this embodiment is formed from a liquid phase of a first preparation liquid and takes a form of fine particles from the action of the water-soluble polymer. The first preparation liquid before precipitation of the water-soluble salt is prepared by a known process, and the water-soluble, other water-soluble salt may be formulated in any order. When the water-insoluble substance is formulated, the water-insoluble substance may be formulated before precipitation of the water-soluble sat dissolved in the first preparation liquid. Caused by precipitation of the water-soluble salt dissolved in the first preparation liquid caused by precipitation of the water-soluble salt, and the water-insoluble substance may be formulated after the precipitation, from the viewpoint of increasing the production efficiency of the second preparation liquid.

[0092] Examples of precipitating a water-soluble salt dissolved in the first preparation liquid will be described below.

5 7-1-1. Precipitation by Addition of Microcrystal-Precipitating Agent

10

30

10093] The process for precipitating the water-soluble salt mentioned above has been studied. As a result, a process of precipitation by means of a microcrystal-precipitation gaent has been found. Specifically, by adding to the first preparation liquid a microcrystal-precipitating agent having an effect of precipitating microcrystals derived from a fine water-soluble salt, the water-soluble salt dissolved in the first preparation liquid before adding the microcrystal-precipitating agent is allowed to precipitate as microcrystals, whereby a second preparation liquid can be obtained. The microcrystal-precipitating agent of the present invention will be described in further detail. Here, from the viewpoint of forming effective supporting sites in the particles for supports, it is preferable that the precipitated water-soluble salt comprises sodium carbonate and/or sodium sulfate.

25 [0094] The microcrystal-precipitating agent refers to a substance which has an effect of precipitating a substance derived from a water-soluble salt different from the precipitating agent by addition to the first preparation liquid.

10095 First, in a case where the microcrystal-precipitating agent is a water-soluble substance, an embodiment where a first preparation liquid comprises a water-soluble sait a and a water-soluble sait before he step of adding a micro-crystal-precipitating agent is described. In this embodiment, the microcrystal-precipitating agent is a substance having a dissolving strength or the water-soluble sait a and the water-soluble sait b at a temperature in which the precipitating agent is added. The term "dissolving strength" as referred herein means extend of easiness in dissolving. The microcrystal-precipitating agent can be variously selected depending upon the kinds of the water-soluble sait contained in the first preparation liquid. A substance which can be used as a microcrystal-precipitating agent can be obtained by the following method. For instance, when a water-soluble substance is added a saturated solution containing the water-soluble sait and the water-soluble sait b, in an embodiment where the objection of the water-soluble sait b, and the water-soluble sait b, in an embodiment where the objection of the water-soluble sait b, in an embodiment where the objection of the water-soluble sait b, in an embodiment where the objection of the water-soluble sait b, in an embodiment where the objection of the water-soluble sait b, in an embodiment where the objection of the water-soluble sait b, in an embodiment where the objection of the objection

dissolved and a substance derived from b, such as b and/or a compound salt or complex salt of a and b, is precipitated, it means that c has a dissolving strength greater than that of b, so that c acts as a microcrystal-precipitating agent. [0096] For instance, when sodium sulfate, sodium carbonate and sodium chloride are added in that order, since sodium chloride is dissolved in a saturated solution of sodium sulfate and sodium carbonate, fine acciual crystals of burkelte, which is a compound salt of sodium sulfate and sodium carbonate, are precipitated without being acgregated.

burkeite, which is a compound salt of sodium sulfate and sodium carbonate, are precipitated without being aggregated. In this case, sodium chloride is a preferable microcrystal-precipitating agent against the preparation liquid comprising sodium carbonate and sodium sulfate.

[0097] The crystals precipitating in the preparation liquid by the microcrystal-precipitating agent are very fine. The size of the crystals precipitating in the second preparation liquid can be determined by using the in-line type powder droplet monitoring system (manufactured by LASENTEC, "TSUB-TEC M100") mentioned above.

[0098] In addition, the effect of pracipitating microcrystals by the microcrystal-pracipitating agent can be confirmed as an increase in the number of particles with the passage of time which is observed after addition of the precipitating agent by the in-line type powder droplet monitoring system.

[0099] As described above, the confirmation of the microcrystal-precipitating agent can be made also in the preparation liquid of any composition, and a method for confirming a microcrystal-precipitating agent in a preparation liquid containing sodium carbonate and sodium suffate will be exemplified.

[0100] First, a saturated solution containing both sodium sulfate and sodium carbonate is prepared by the following method. Four-hundred grams of sodium sulfate (curity 99% or more) is added to 1500 g of fon-exchanged water, which is adjusted to the preparation temperature of the first preparation liquid. The mixture is sufficiently stirred to 20 minutes in a thermostat set at the preparation temperature of the first preparation figuid to dissolve sodium sulfate. Further, 400 g of sodium carbonate (TDENSE ASHT manufactured by Central Glass Co., Ltd.) is added thereto, and the mixture is stirred for 30 minutes, to give a suspension. A saturation solution of sodium sulfate/sodium carbonate is prepared by a method of following superantiant after allowing the suspension to stand, or by a method of firsting the suspension.

Here, the term "the preparation temperature of the first preparation liquid" refers to any temperature within the temperature range of from 30° to 80°C.

[0101] One-thousand grams of the saturated solution of sodium sulfate/sodium carbonate prepared in the manner described above is weighed and pisod in a 1-L stainless beaker, and stirred in a thermostat of which temperature is adjusted to the same temperature as that of the preparation liquid with rotating apitation impellers with 3 propeller wings of 2 × 4 cm at a speed of 200 r/min. The measurement is initiated in the same namera as described above by using the in-line type powder droplet monitoring system, manufactured by LASENTEC. A 100 g test sample is added within 30 seconds, and proceeded with 60-minute stirring and measurement. When at least any one of microcrystals derived from sodium carbonate and/or sodium suitate, of which average particle size after 90 minutes (code length at which the cumulative value of the number of particles is 50%) is 40 µm or less, for instance, sodium carbonate and dyrdrates thereof, sodium suitate, and short as the sodium suitate, is precipitated, the test sample is a microcrystal-precipitating agent against sodium carbonate and sodium suitate, in addition, the average particle size of the precipitated microcrystals is more preferably 30 µm or less, film ore preferably 20 µm or less, most preferably 10 µm or less. Here, the precipitate is identified by analyzing with X-ray diffraction, elemental analysis, and the like.

[0102] The microcrystal-precipitating agent includes, for instance, salts having high dissolving strength such as chicnides, bromides, iodides and fluorides of alkali metals and/or alkaline earth metals, such as sodium, potassium, calcium and magnesium. In addition, there may be also included, as microcrystal-precipitating agent, sovents which are compatible with water such as ethanol, methanol, and acetone; and substances having a large hydration force, such as scellate (anhydride). In other works, by dissolution, hydration, and the like of the microcrystal-precipitating agent, water used for dissolution of the water-soluble salt in the first preparation liquid is taken away, thereby serving as a base material having an effect of precipitating the water-soluble salt from the liquid phase of the first preparation liquid. [0103] From the viewpoint of the dissolving strength, the bromides and lodides are preferable, and from the viewpoint of the storage stability of the detergent particles, the chlorides are preferable. Also, from the viewpoint of the influence, the alkali metals alsts are preferable. Also, from the microm the economic, viewpoint

sodium chloride is especially preferable.

[1014] The contrator of the microcrystal-precipitating agent in the particles for supporting a surfactant is preferably

from 0.2 to 35% by weight, more preferably from 0.5 to 30% by weight, more still preferably from 1 to 25% by weight, more

particularly preferably from 2 to 20% by weight, sepcially preferably from 4 to 15% by weight, from

exhibiting a sufficient effect for microcrystal precipitation and the viewpoint of maintaining the detergency performance

when used as a deterenat composition.

25

50

[0105] In addition, it is preferable that as to the dissolution rate of the water-soluble, microsystal-precipitating agent in the second preparation liquid, the higher the dissolution rate, the better, from the viewpoints of generating a large amount of precipitates in the second preparation liquid by largely dissolving in the solution portion of the first preparation upon the precipitate in the second preparation liquid by largely dissolving in the solution portion of the first preparation liquid, so as to have a preferable structure for a supporting site in the particles for supports obtainable after spray-dying to the filling drompostion. The dissolution rate of the microsystal-precipitating agent is preferably 75% by weight or more, more preferably 85% by weight or more, most preferably 85% by weight or more, most preferably being completely dissolved.

[1016] The dissolution rate of the microcystal-precipitating agent in the second preparation liquid can be determined by combining known analyzing means. For instance, the second preparation liquid is filtered under reduced pressure, and thereafter the water concentration P (%) in the filtrate is measured with a far infrared ray heather-type moisture meter (manufactured by SHMAZD CORPORATION) or the filtrate. Further, the concentration of the microcystal-precipitating agents (%) in the filtrate is obtained by ion chromatography or the like. Supposing that the water content of the second preparation liquid is T (%), the dissolution rate of the microcystal-precipitating agent in the second preparation records to the proper second preparation records of the proper second preparation records and the property second preparation records and

Dissolution Rate (%) =
$$\frac{(100 \times S \times Q)}{(P \times T)}$$
 (VI)

[0107] In an embodiment where sodium carbonate and sodium sulfate are together contained in the first preparation liquid, it is preferable that sodium carbonate is added after sufficiently dissolving sodium sulfate, from the viewpoint of increasing the supporting ability of the particles for supports.

[0108] The water content of the second preparation liquid is preferably from 30 to 70% by weight, more preferably from 35 to 56% by weight, most perferably from 35 to 56% by weight, most perferably from 40 to 60% by weight, from the viewpoints of reducing undiscoved substances of the water-soluble components which are not microcrystals and effectively exhibiting the effect of the microcrystal-precipitating agent. The temperature of the preparation liquid be preferably from 30° to 80°C, more preferably from 30° to 75°C, from the viewpoints of the dissolved amount of the water-soluble sait and the liquid convey.

ability with a pump.

[0183] Concrete examples of the process for preparation of this embodiment include, for instance, initially adding all or substantially all of water to a mixing vessel, and sequentially adding other components, preferably after the water temperature almost reaches a set temperature, to give a first preparation liquid. A preferable order of addition is such that liquid components and sodium suffice, sodium carbonate, and the like are initially added. In addition, small amounts of auxiliary components such as water-insoluble substances, such as zeolite, and dyse can be also added. The incrocrystal-precipitating agent is added in a state where the solution portion of the first preparation liquid is saturated. Alternatively, in a case where the solution portion is in an unsaturated state, the microcrystal-precipitating agent is added in an amount exceeding that necessary for the solution portion to be saturated. The water-insoluble substance may be added before the addition, after the addition or in divided portions before and after the addition of the microcrystal-precipitating agent. In order to finally obtain a homogeneous second preparation liquid, after the addition the entire components to the preparation liquid, the mixture is mixed for preferably 10 minutes or more, more preferably 30 minutes or more.

7-1-2. Precipitation by Concentration of First Preparation Liquid

[0110]. The process for precipitating the water-soluble salt mentioned above has been studied. As a result, a process of precipitating by concentrating the preparation liquid has been found. In other words, a large number of microcrystats can be generated in the second preparation liquid by carrying out the operation of precipitation by means of concentration of the water-soluble salt in a dissolving state in the presence of the water-soluble polymer. The concentration of the preparation liquid in this embodiment will be described in further detail.

[0111] A process of obtaining a concentrated sturry in which a part of the water-soluble salt dissolved in the first preparation liquid is precipitated by concentrating the first preparation liquid comprising a water-soluble polymer and a water-soluble salt will be described.

[0112] First, the first preparation liquid before concentration may be prepared by a known process, and the watersoluble polymer and the water-soluble salt may be formulated in any order. In addition, in a case where a water-installed substance is formulated, the water-insoluble substance may be formulated before concentration of the first preparation liquid, or it may be formulated afterwards. In addition, the concentration operation may be carried out to the second preparation liquid subjected to a treatment, for instance, formulation of a microrystal-precipitating agent or the like.

[0113] The smaller the amount of the coarse particles of the undissolved water-soluble salt which are present in the first preparation liquid before concentration, the higher the supporting ability of the particles for supports obtained the spray-drying. Therefore, the dissolution rate of the water-soluble salt in the first preparation liquid before the concentration is preferably from 50 to 100% by weight, more preferably from 70 to 100% by weight, especially preferably from 80 to 100% by weight. When the dissolution rate does not reach 100% by weight, there is a preferable embodiment where the undissolved substances are made finer by pulverizing the first preparation liquid by using the subsequently described wet pulverization device or the like. The wet pulverization of the first preparation liquid may be carried out for 5 concentrated slurry. Here, the dissolution rate of the water-soluble salt is determined by the method described above.

[0114] Next, the water-soluble salt dissolved in the first preparation liquid is precipitated by concentrating the first preparation liquid. The extent of the concentration can be determined by the amount of water loss in the first preparation liquid. The amount of water loss in the first preparation liquid is not particularly limited, and it may be appropriately set so that the amount of the water-soluble salt precipitated takes a preferred range. In addition, the water content in the first preparation liquid is not particularly limited. As a device for concentration, it may be any sorts of concentrators which are widely used. For instance, a natural circulation evaporator in which a liquid naturally dirculates by rising with boiling within a heating tube; in forced circulation evaporator with external heating in which a liquid is circulated at a high speed between an evaporator and a heater with a circulation pump, and water is evaporated with an evaporator; and a falling thin-film evaporator in which a liquid is allowed to flow into the evaporator from a top of a vertical heater, and subjected to evaporation and concentration by forming homogeneous liquid film on the inner wall of the heater during a falling. These evaporated by ejecting a liquid heated to a temperature of a boiling point or higher in the evaporator under reduced pressure is also effective.

[0115] Since the first preparation liquid used in this embodiment produces crystals of the water-soluble salt along with the concentration, the scales are likely to deposit in the concentrator. Therefore, it is more preferable to use a concentrator having a function that can remove the deposited scales, or a concentrator having a structure in which scales are less likely to deposit. A former device includes a device in which the above-described falling thin-film evaporator is equipped with agitation impellers for scraping off the scales, for instance, Wipten (manufactured by Shinkton Pantec Co., Ltd.). A latter device includes losoo evaporator (manufactured by SUMITOMO HEAVY INDUSTRIES,

LTD.) which comprises a plate-type heating element in the inner portion thereof, in which concentration is carried out by allowing to flow a liquid on a surface of this heating element under reduced pressure.

7-1-3. Precipitation by Reduction of Dissolved Amount by Temperature Adjustment of First Preparation Liquid

6

[0116] The process for precipitating the water-soluble salt mentioned above has been studied. As a result, a process of precipitating the water-soluble salt by changing the temperature of the first preparation liquid so as to lower the dissolved amount of the water-soluble salt has been found. In other words, a large number of microcrystals can be precipitated in the preparation liquid by emperature-adjustment so as to lower the dissolventying amount of the water-soluble salt in a dissolved state in the first preparation liquid, thereby allowing to precipitate the water-soluble salt in a dissolved state in the presence of a water-soluble polymer. The precipitation by reduction of the dissolved amount by temperature-adjustment of the preparation liquid in this embodiement will be described in further destination.

[0117] A process of obtaining a second preparation liquid in which a part of the dissolved water-soluble salt is precipitated by changing the temperature of a first preparation liquid so as to lower the dissolved amount of the watersoluble salt in the first preparation liquid comprising a water-soluble polymer and a water-soluble salt will be described. [0118] First, the first preparation liquid comprising a water-soluble polymer and a water-soluble salt will be described. [0118] First, the first preparation liquid comprising a water-soluble solution may be prepared by a known process. Also, when a water-insoluble substance is formulated, the water-insoluble substance may be formulated before the temperature-changing operation of the first preparation liquid, or after the emperature-changing operation. In addition, a part of the water-soluble polymer may be formulated in the second preparation liquid after the above operation. By the formulation, the size of the precipitated water-soluble salt crystals can be also adjusted, Also, the temperature-changing operation may be carried out to the second preparation liquid in the same manner as in the concentration oneration.

[0119] The smaller the amount of the coarse particles of the undissolved water-soluble salt which are present in the first preparation liquid before the temperature-changing operation, the higher the supporting a sulfactant. Therefore, the dissolution rate of the water-soluble salt in the first preparation liquid before the temperature-changing operation is referably from 50 to 100% by weight, especially preferably from 90 to 100% by weight, the proclaip type-derably from 90 to 100% by weight, there is a preferable embodiment where the undissolved substances are made finer by pulverizing the first preparation liquid by using the subsequently described wet pulverization device or the like. The well pulverization device or the like. The well pulverization of the preparation liquid may be carried out to the second preparation liquid after the temperature-changing operation. Here, the dissolution rate of the water-soluble salt is determined by the method rescribed about 1900 frescribed and the subscribe water soluble salt is determined by the method rescribed about 1900 frescribed about

[0120]. Next, a part of the dissolved water-soluble salt is precipitated by changing the temperature of the first perparation liquid. A process of changing the temperature of the first preparation liquid includes a process of heating or cooling the first preparation liquid by using a device equipped with an external jacket, an internal coil, or the like when preparing the first preparation liquid, or the like.

[0121] It is preferable that the temperature of the first preparation liquid before the temperature-changing operation is set so that the dissolution rate of the water-soluble sait contained in the preparation liquid is high, and an optimal temperature is determined by the kinds and the amounts of the formulated water-soluble sait.

[0122] The temperature of the second preparation liquid after the temperature-changing operation is set so that the dissolution rate of the water-soluble sait in the preparation liquid is lowered, and selection of heating or cooling must be made depending upon the kinds and amounts of the formulated water-soluble salt. Sodium sulfate and sodium carbonate sultably used as detergent raw materials show a maximum dissolved amount near 40°C. Therefore, when these raw materials are used, its preferable that the temperature of the first preparation liquid before the tomperature-changing operation is adjusted to 40°C or so, and that the temperature of the second preparation liquid after the temperature-changing operation is adjusted to 50° to 70°C.

[0123] Here, there is also a preferable embodiment in which the precipitation of the dissolved water-soluble salt is accelerated, for example, by subjecting the preparation liquid to flash concentration together with changing the temperature of the first preparation liquid.

7-2. Wet Pulverization of Water-Soluble Salt Particles in First Preparation Liquid

[0124] This embodiment comprises (a) preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble polymer and a water-soluble salt; and (b) subjecting water-soluble salt particles in the first preparation liquid to wet pulverization. In this embodiment, the first preparation liquid before wet pulverization may be prepared by a known process, and the water-soluble polymer and the water-soluble salt may be added in any order. A water-insoluble substance may be formulated before subjecting the first preparation liquid to wet pulverization, or it may be formulated after subjecting the first preparation liquid to wet pulverization. From the viewpoint of uniform dispersion by disinterartian the accreaated mass of the water-insoluble substance. It is referable to formulate before the wet qui-

- verization treatment. For instance, the calcium exchange speed of the crystalline aluminosilicate can be improved. [0125] In addition, the finer the water-soluble salt particles which are present in in the first preparation liquid are pulverized, the larger the effect of improving the supporting ability of the particles for supporting a surfactant obtainable in the subsequent sore-deviation process.
- 5 [0126] The wet pulverization treatment of the first preparation liquid can utilize the water-soluble salt in the formation of supporting sites in the particles for supports obtainable in the subsequent spray-drying process by pulverizing the water-soluble salt particles in the preparation liquid. The wet pulverization has an especially large effect when a water-soluble solid derived from soldium carbonate is present in the first preparation liquid. As a concrete illustration, when burketie, which is a compound salt derived from soldium carbonate, is formed in the first preparation liquid betained by the indirection of the preparation liquid coaling the properties of the preparation liquid betained by the indirection of the supporting sites in the particles for supports when being present as coarse particles, it can be effectively utilized in the formation of the supporting sites in the particles for supports by making them fine by wet pulverization, whereby the supporting ability of the particles is improved.
- 15 [0127] In addition, the advantages of a case where sodium carbonate is formulated in the first preparation liquid are as follows. In an embodiment where sodium carbonate is formulated in the preparation liquid by finely pulverizing by a dry-type pulverizer, the undissolved substances undesitably form coarse particles by aggregation upon hydration. However, in an embodiment where sodium carbonate is formulated in the first preparation liquid and thereafter the mixture is subjected to wet pulverization, the formation of the coarse particles by the aggregation described above can be suppressed.
 - [0128] Conditions for the wet pulverization treatment cannot be limited absolutely, as they depend on the concentration of the water-soluble sait in the first preparation liquid, the pulverizers used, and the like. The pulverizers which can be used in this embodiment may be any ones, as long as they are generally known were pulverizers. The usually employed wet grinders include (i) devices in which fine pulverization is carried out by utilizing pulverization media; and (ii) devices in which fine pulverization tracifed out with a gap between a pulverization side and a stator.
 - [0129] The device (i) includes a device in which pulverization is carried out with a shearing force caused by the difference between the flow rates of the media by supplying a solution to be treated from the bottom of the vessel, and discharging the solution to be treated from the top of the vessel, with sifring the media inside the vertical cylindrical vessel with sighted in pulperial solution in pellers and an agitation disc. Such continuous process-type devices include a sand grinder (manufactured by ligaresh Kika Selzo K.K.), and a universal mill (manufactured by K.K. Missu Milke Seisakusho); and batch process-type devices include ACUJAMIZER (manufactured by Hosokawa Micron Corporation). Horizontal continuous process-type devices having a similar structure include DYNOMILL (manufactured by MAB). Also included are those comprising a cylindrical rotor and an annular casing enveloping it, in which pulverization of a solution to be treated fed from the bottom center of a rotor is carried out by high-speed-rotation force of the media, including DIAMOD FINE.

 MILL (manufactured by Mitsubishi Heavy Industries, Ltd.); and KOBOL MILL (manufactured by Shinko Pantec Co., Ltd.).
 - [0130] The device (ii) includes those comprising a rotor and a stator each having grinding (seth, in which putverization is carried out by repeatedly applying a shearing force when the solution to be treated is passed through the gap, including Collold Mill (manufactured by Shinko Pantec Co., Ltd.), and Trigonal (manufactured by Missu Milke Machinery Co., Ltd.). Along the activated as the saving a similar grinding mechanism, except that a rotor and a stator is a grinding store, including Glo-Mill (manufactured by K.K. Glo Engineering), Super Maskoroider (manufactured by Masuko Sangyo K.K.), and Corandom Mill (manufactured by Shinko Pantec Co., Ltd.). Also included is one in which that solution to be rested is roughly pulverized with a first turbine and a stator, and the roughly pulverized mixture is then finely pulverized with a second rotor and a stator, inducing Homomix Line Mill (manufactured by Toxibu Kika Kogyo K.K.). Further included is one in which a dispersion effect of the level of high-pressure homogenizer can be attained by applying to the liquid a strong impact of the order of megahertz with a wet-type emulsification disperser having all of the functions of emulsification and dispersion, homogenous mixing, and finely providering by a rotator having a peculiar shape and being high-speed rotated and a stator which is engaged therewith, including CABITRON (manufactured by PACIFIC MACHINERY & ENISINEERING Co., Ltd.).

7-3. Addition of Fine Particles to Preparation Liquid

50

[0131] This embodiment comprises (a) preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble path, and (b) adding to the first preparation liquid fine water-soluble salt particles are capable of being present without substantially being dissolved in the first preparation liquid. In this embodiment, the phrase "under the conditions that fine water-soluble salt particles are capable of being present without substantially being dissolved in the first preparation liquid." means that when the solution portion of the first preparation liquid is saturated, the added fine particles are not dissolved.

and that when the solution portion is in an unsaturated state, the fine particles dissolve until the solution is saturated by the addition thereof, but once the saturation is reached, no more fine particles are dissolved. The fine water-soluble salt particles are those salts which are substantially the same as the water-soluble salt which remains undissolved in the first preparation liquid and/or the same sait as the water-soluble salt firstly precipitated and/or those salts having the smallest dissolving strength in the second preparation liquid.

[0132] In addition, the first preparation liquid berofe addiring the fine water-soluble salt particles is prepared by a known process, the first preparation injuried before the first preparation profit. When the water-insoluble substance is formulated in any other profit p

[0133] Here, as the fine water-soluble salt particles mentioned above, the fine particles having substantially the same composition as the firstly precipitated water-soluble salt from the first preparation liquid are preferable. The phrase "the fine particles having substantially the same composition as the firstly precipitated water-soluble salt from the first preparation liquid" refers to fine particles having substantially the same composition as a substance precipitated when a part of moisture in the first preparation liquid before adding the fine particles is evaporated, and/or as a substance precipitated when the temperature is changed, when the water-soluble salt particles are not present in the first preparation liquid before addition of the fine particles. Here, as a process for preparing the fine particles, there can be considered to fine pulverization of the commercially available appropriate substances, and it is more preferable to form microcrystals in the presence of the water-soluble polymer. Concretely, a substance having the same composition as the fine particles is dissolved in water together with the water-soluble polymer, and is allowed to crystallize by spraydrying or the like, and the crystals are made fine with a pulverizer, to give fine particles. The fine pulverizers include roller mills, ball-mills, collision-type pulverizers, and the like. The roller mills include USV mill (manufactured by Ube Industries, Ltd.), MRS mill (manufactured by Mitsubishi Heavy Industries, Ltd.), SH mill (manufactured by IHI), and the like; the ball-mills include Dynamic Mill (manufactured by Mitsul Milke Machinery Co., Ltd.), Vibration Mill (manufactured by Chuo Kakoki Shoji K.K.), and the like; and the collision-type pulverizers include Atomizer, Pulverizer (both being manufactured by Fuji Paudal Co., Ltd.), and the like.

[0134] In addition, the smaller the average particle size of the fine particles, the larger the effect of improving the supporting ability of the particles for supporting a surfactant obtainable by spray-dying in the subsequent process. From this viewpoint, the average particle size of the fine particles is preferably 40 µm or less, more preferably 35 µm or less, still more preferably 30 µm or less, still more preferably 25 µm or less, still more preferably 10 µm or less. Here, the average particle size is determined by the following method.

[0135] One-thousand grams of ethano is weighed and placed into a 1-L stainless beaker, and stirred in a thermostat at 20°C with rotating apitation impeliers with 5 propeller wings of 2 × 4 cm at a speed of 200 rimin. Subsequently 20 g of the fine particles mentioned above are supplied. The particle size distribution at a point of measuring for 10 minutes is determined in the same manner as described above by using the in-fine type powder droplet monitoring system, manufactured by LASENTEC (TSUB-TEC M100). Here, a median code (particle size at which the cumulative value of the number of particles is 20%) is considered as an average particle size.

[0136] In addition, in the embodiments described above, regarding step (b), it is preferable that the treatment of increasing the number of the water-soluble sail particles comprises one or more processes selected from the group consisting of (1) adding a microcrystal-precipitating agent to the first preparation liquid; (2) concentrating the first preparation liquid; (3) adjusting a temperature of the first preparation inquid so that the dissolved amount of the water-soluble sail to lowered; (4) subjecting water-soluble salt particles in the first preparation inquid to wet pulverization; and (5) adding to the first preparation liquid fine water-solubles and the process of different from the water-soluble salt in the first preparation liquid, under conditions that the fine water-soluble salt particles are capable of being prepent without substantially being dissolved in the first preparation liquid.

[0137] By carrying out the steps (a) and (b) in the embodiments as described above, the second preparation liquid is obtained.

8. Process for Preparing Cave-In Particle

50

[0138] It is preferable that in the particles for supports of the present invention, at least a part of particles is composed of a particle which is a cave-in particle having a structure that there exists a hollow, namely a cave-in hole, in an inner portion thereof, and that a particle surface is opened and communicated with the hollow in the inner portion. The particles for supports are prepared by providing holes with a very line needle and the like from the surface to the inner portion of the particle to which a surfactant can be supported.

[0139] In addition, a process for effectively preparing the cave-in particle in the present invention includes a process comprising adjusting a surfactant content of the second preparation liquid mainly comprising a water-soluble and to obtained in the manner described above to from 0 to 2% by weight, and adjusting a water soluble salt obtained in the manner described above to from 0 to 2% by weight, and adjusting a water soluble salt obtained in the manner described above to from 0 to 2% by weight, and adjusting a water soluble salt obtained in the manner described above to from 0 to 2% by weight, and adjusting the complete salt of the control of the contro

content of the second preparation liquid having an increased number of water-soluble salt particles to a range of from 35 to 65% by weight, and spray-drying the preparation liquid.

[0140] In the present invention, there is exhibited an effect that the content of the cave-in particle in the spray-dried particle is remarkably increased by adjusting the surfactant content and the water content of the second preparation figured to the ranges as specified above, respectively, and increasing the number of the water-soluble salt particles in the second preparation figurif, namely by allowing the water-soluble salt to be present in an undissolved state.

[0141] The content of the surfactant in the second preparation liquid is from 0 to 2% by weight, preferably from 0 to 1% by weight, more preferably 0% by weight, from the viewpoint of increasing the content of the cave-in particle in the particle so total nable by spray-drying the preparation liquid.

[0142] The water content of the second preparation liquid is preferably from 35 to 65% by weight. In addition, the water content is 35% by weight or more, preferably 37% by weight or more, preferably 39% by weight or more, still more preferably 41% by weight or more, most preferably 45% by weight or more, most preferably 45% by weight or more, from the viewpoints of making the supporting capacity of the particles for supports larger and opening a cave-in hole of a sufficient size. Also, the water content is 65% by weight or less, preferably 62.5% by weight or less, more preferably 60% by weight or less, such supports are preferably 55% by weight or less, more preferably 60% by weight or less, from the viewpoint of suppressing the bursting of the droplets by the temperature elevations.

[0143] In addition, as contents of other components in the second preparation liquid, the water-soluble polymer is contained in an amount of preferably from 1 to 20% by weight, more preferably from 3 to 15% by weight, still more preferably from 5 to 10% by weight, the water-soluble salt is contained in an amount of preferably from 7 to 59 by weight, more preferably from 20 to 35% by weight, Further, when the water-insoluble substance is contained; the an amount of preferably from 3 to 32% by weight, more preferably from 10 to 18% by weight.

[0144] . The preparation liquid having the composition described above may be those which are liquid-conveyable and non-curable. In addition, the addition method for each component and its order can be appropriately varied depending upon the conditions.

[0145] In addition, in the second preparation liquid, a part of the water-soluble salt is present in an undissolved state. In the present invention, there are advantages in the preparation liquid as described above in that a cave-in hole is generated in the particles for supports and a supporting ability for the liquid surfactant composition can be enhanced by allowing a part of the water-soluble salt to be present in an undissolved state.

0 [0146] The undissolved amount of the water-soluble salt is preferably from 0.5 to 15% by weight, more preferably from 1 to 11% by weight, still more preferably from 2 to 9% by weight, most preferably from 3 to 7% by weight, of the second preparation liquid. In addition, the undissolved water-soluble salt particles mentioned above (hereinafter also referred to as "undissolved substance") have an average particle size of preferably 80 µm or less, soll more preferably 40 µm or less, sepocially preferably 80 µm or less.

[0147] Here, a process for allowing the undissolved substance to be present in the second preparation liquid includes, for instance, a means of adjusting the content of the water-soluble salt and the content of water to those with the ranges described above, a means of adjusting a temperature of the preparation liquid by considering the dissolved amount of the water-soluble salt, and the like, in addition, the means of making a particle sate of the undissolved substance smaller includes means described above such as a means of adding fine water-soluble salt particles to a first preparation liquid, under conditions that the fine particles are capable of being present without substantially being dissolved in the first preparation liquid, a means of making its size smaller by a means of purvisiting the retired of the first preparation liquid, thereby precipitating the crystals; a means of sovering a dissolved amount by varying a temperature of the first preparation liquid, thereby precipitating the crystals; a means of soverparating an incrocrystal-precipitating agent to the first preparation liquid, thereby precipitating encystals of the water-soluble salt which is dissolved therein, and the like. [0148] Here, with regard to the determination of the undissolved amount of the water-soluble salt, the second preparation liquid, thereby collecting supernation, manyly the solution portion of the second preparation liquid is centrifuged, thereby collecting supernation, manyly the solution portion of the second preparation liquid centrifuged.

About 3 g of the solution is weighed with an accurate balance in an amount of a (g), and dried at 165°C for 4 hours. Thereafter, the resulting solution is cooled in a deslicator for 30 minutes, and the dried remnant of the supernatant is weighed with an accurate balance in an amount of b (g). Here, the dissolved amount of the supernatant c (%) is

calculated by:

50

20

$$\frac{b}{a-b} \times 100$$

Also, the content d (%) of the water-soluble salt contained in the dried remnant is analyzed. Using the water content e (%) of the second preparation liquid and the content f (%) of the water-soluble salt in the second preparation liquid, the undissolved amount (%) of the water-soluble salt is calculated by the following equation:

Undissolved Amount (%) = f-e
$$\times \frac{c}{100} \times \frac{d}{100}$$
 (VII)

[0149] In addition, with regard to the measurement of the average particle size of the undissolved water-soluble salt, the average particle size can be determined by using the in-line type powder droplet monitoring system (manufactured by LASENTEC, "ISUB-TEC M100") mentioned above.

[0150] The second preparation liquid is obtained by obtaining a first preparation liquid by a known process, and thereafter subjecting the preparation liquid to a treatment of increasing the number of the water-soluble salt particles mentioned above.

0 [1511] In the spray-drying process, a method for generating a cave-in particle in the particles for supports includes, though differences are caused in the original control ranges by the difference in the composition for the particles for supports, a means of controlling to a range of drying conditions suitable for the composition, and a means for controlling the water content of the second preparation liquid.

[0152] In the control for the drying conditions, it is preferable that conditions which quickly dry the sprayed droplets, anamely a temperature of the periphery of the droplets immediately after spraying is preferably 85°C or more, more preferably 90°C or more, However, from the viewpoint of thermal degradation of the constituents, the air blow temperature is preferably 400°C or less, more preferably 350°C or less, secolally preferably 300°C or less.

20 9, Properties of Particles for Supporting Surfactant

[0153] The bulk density of the particles for supports of the present invention is preferably from 300 to 1000 glt. more preferably from 450 to 80 gl, still more preferably from 450 to 80 glt., from

[0154] In addition, from the viewpoints of generation of fine powder dusts and dissolubility when using a detergent composition comprising detergent particles comprising particles for supports and a liquid surfactant composition supported thereby, the average particle size of the particles for supports is preferably from 140 to 600 µm, more preferably from 160 to 500 µm, still more preferably from 180 to 400 µm.

0 [0155] The supporting capacity for a preferable liquid surfactant composition to the particles for supports is 0.35 mL/g or more, more preferably 0.45 mL/g or more, especially preferably 0.56 mL/g or more, most preferably 0.50 mL/g or more, from the viewpoint of increasing the permitted range of the formulation amount of the liquid surfactant composition.

[0156] A preferable supporting rate of the particles for supports is preferably 0.2 mL/g or more, more preferably 0.3 at mL/g or more, sill more preferably 0.4 mL/g or more, from the viewpoint of more quickly and efficiently absorbing the liquid surfactant composition, thereby increasing the productivity.

[0157] The lower the water content of the particles for supports as determined by an infrared moisture meter, the better, from the velopoint of making the supporting capacity for the fujuid surfactant composition of the particle starger. The water content is preferably 14% by weight or less, more preferably 10% by weight or less, still more preferably 5% by weight or less.

[0158] Here, the bulk density, the average particle size, the supporting capacity for the liquid surfactant composition, the supporting rate, and the water content can be determined by the method described under the method for determining properties described below.

45 10. Composition and Properties of Detergent Particles

[0159] The detergent particles of the present invention comprise the surfactant for supports mentioned above and a surfactant composition supported therein.

[0160] In the surfactant composition, an anionic surfactant and a nonionic surfactant can be each used alone, and it is more preferable to use both surfactants in admixture. Especially in a case of using a nonionic surfactant having a melting point of 30°C or less, it is preferable to use it in combination with a water-soluble nonionic organic compound (hereinafter referred to as "melting point-elevating agent") having a melting point of from 45° to 100°C and a molecular weight of from 1000 to 3000, or an aqueous solution thereof, which has a function of elevating a melting point of this nonionic surfactant. Here, the melting point-elevating agent which can be used in the present invention includes, for instance, polyethylene glycols, polypropylene glycols, polyoxyethylene alkyl ethers, pluronic type nonionic surfactants, and the like. In addition, an amphoteric surfactant or a cationic surfactant can be also used in combination therewith in accordance with its purpose. Also, since an anionic surfactant such as an alkylbenzenesulfonate is formulated in the detergent particles in an amount of from 5 to 25% by weight, an effect of improving the dispersibility of the detergent

particles in low-temperature water is exhibited.

25

30

[0161] As the surfactant composition, there can be used, for instance, one or more kinds selected from the group consisting of anionic surfactants, nonionic surfactants, cationic surfactants and amphoteric surfactants. The anionic surfactants are exemptified by alkybenzenessulfonates; alky either or alkenyl ether sulfates; co-befinsulfonates; salts of α-sulfonated fatty acids or esters thereof; alkyl ether or alkenyl ether carboxylates, amino acid-type surfactants; N-acyl amino acid-type surfactants, and the like. Especially included are linear alkybenzenesulfonates of which alkyl moiety has 10 to 14 carbon atoms; and alkyl sulfates or alkyl ether sulfates, of which each alkyl moiety has 10 to 18 carbon atoms. The counter ions are preferably alkali metals such as sodium and potassium, and amines such as monoethanoidamine and diethnolamine.

[0162] Further, in order to obtain defoaming effects, a fatty acid salt can be used in combination therewith. The preferable number of carbon atoms of the fatty acid molety is from 12 to 18.

[0183] The nonionic surfactants include polyoxyethylene alkyl or alkenyl ethers, polyoxyethylene alkyl or alkenyl ethers, polyoxyethylene-polyoxypropytene alkyl or alkenyl ethers, polyoxyethylene-polyoxypropytene glycols are presented by the trade name "pluronic", polyoxyethylene elkylamines, higher fatty acid alkanolamides, alkyl glucosides, alkyl glu

R-O-(EO)_Y-(PO)_Y-(EO)₇-H

wherein R is a hydrocarbon group, preferably an alkyl group or an alkenyl group; EO is an oxyethylene group; PO is an oxypropylene group; and X, Y and Z are each average moles thereof, among which most preferable average moles have the relations of X > 0.7 > 0.1 × Y+Z = 6 to 14; X+Z = 5 to 12; and Y = 1 to 4.

[0164] The cationic surfactants include quaternary ammonium salts such as alkyl trimethyl ammonium salts.

[0165] The amphoteric surfactants are exemplified by carbobetain-type and sulfobetain-type surfactants and the like. [0166] The formulation amount of the anionic surfactant is preferably from 0.0 300 parts by weight, more preferably from 20 to 200 parts by weight, especially preferably from 30 to 180 parts by weight, based on 100 parts by weight of the nonlonic surfactant. The formulation amount of the melting point-elevating agent of the nonlonic surfactant is preferably from 1 to 100 parts by weight, more preferably from 1 to 100 parts by weight, more preferably from 1 to 100 parts by weight, more preferably from 1 to 50 parts by weight, based on 100 parts by weight of the nonlonic surfactant. In the above range, the composition is preferable, because the composition has a temperature range so that the viscosity of the composition at a temperature of a pour point or higher is adjusted to 10 Pars or less, preferably 5 Pars or less, especially preferably 2 Pars or less, and also has a temperature range so that the inserting hardness of the composition in the temperature range lower than the pour point of the composition and higher than the melting point of the nonionic surfactant is 10 kPa or more, repreferably 30 Pars or more, especially preferably 50 Pars or more, especially prefera

[0167] The values for the properties of the surfactant composition can be determined by the following method. The pour point can be measured by the method according to JIS K 2289. The metiting point is determined by using FP800 Thermosystem "Mettler FP81" (manufactured by Mettler Instrumente AG) and heating at a heating rate of 0.2°C/min. The viscosity is obtained by measuring with a B-type viscometer ("DVM-B model" manufactured by TOKYO KEIKI),

rotor No. 3 under the condition of 80 r/min. In addition, when the measurement value under the above conditions exceeds 2 Pars, to be undeterminable, the viscosity is obtained by measuring with rotor No. 3, under the condition of 12 r/min. The inserting hardness is a value obtained by determining a load when an adaptor is inserted for 20 mm at an inserting rate of 20 mm/min into an inner portion of the surfactant composition by using a rheometer ("NRM-3002D" manufactured by Fudo Kogyo K.K.) and a disc-shaped adaptor (No. 3, 89) having a diameter of 8 mm and a bottom area of 0.5 cm², and dividing the resulting load by the bottom area of the disc-shaped adaptor.

[0168] The amount of the surfactant composition is preferably in a range of from 10 to 100 parts by weight, more preferably in a range of from 20 to 80 parts by weight, especially preferably in a range of from 30 to 60 parts by weight, based on 100 parts by weight of the particles for supports, from the vlewpoints of the detergency and the dissolubility.

The "amount of the surfactant composition" as referred to herein does not include the amount of the surfactant even if the surfactant were added to the preparation liquid.

[0169] When the surfactant composition is mixed with the particles for supports, powdery raw materials other than the particles may be added as desired, and the amount thereof is preferably from 0 to 150 parts by weight, based on 100 parts by weight of the particles. The powdery raw materials include, for instance, aluminosilicates, crystalline silicates such as SKS-6 (manufactured by Clarient), and the like.

[0170] In addition, the detergent particles can contain the water-soluble polymer, the water-soluble salt, the water insoluble substance, and other components, each of which is exemplified in the particles for supports as components other than the above-mentioned surfactant composition. In a case where a water-insoluble substance is used, the crystalline silicates described below and the like can be also contained.

[0171] Here, when the detergent particles are prepared by using components such as a suffactant which can serve as a binder, and powdey raw materials, the detergent particles are coated with an aggregated layer formed by the above components, so that there may be some cases where the shape of the particles for supports cannot be confirmed simply from their external appearance. A method of differentiating be shape of the particles for supports in such cases includes a method of confirming the shape by extracting an organic solvent-soluble component from the detergent particles, they separating the particles for supports. The kinds of the organic solvent-soluble component from the detergent particles, thereby separating the particles for supports. The kinds of the organic solvents used in extraction are appropriately selected depending upon the kinds of the binder substances bound to each constituent unit of the detergent.

[0172] The method for confirmation of a shape of the particles for supporting a surfactant by solvent extraction will be illustrated hereinbelow.

[0173] Fifteen grams of the detergent particles which are accurately sample-reduced and weighed are subjected to reflux operation for 1 hour with 300 mL of 95% ethanol heated in a water bath. Thereafter, an ethanol-insoluble component is gradually filtered off by means of suction filtration with sufficiently washing with hot ethanol. The separated ethanol-insoluble component is dried for 24 hours under reduced pressure, and thereafter the insoluble component is cautiously collected so as not to disintegrate the particle structure of the insoluble component. Such an operation is carried out several times, to obtain 100 g of an ethanol-insoluble component. The resulting ethanol-insoluble component is vibrated for 10 minutes with standard sieves according to JIS Z 8801. Thereafter, the weight on each sieve is measured, and the particle classified in accordance with each sieve-opening mentioned above is observed and analyzed, to confirm whether or not the resulting particles are the particles for supports of the present invention, or to confirm the absence or presence of an ethanol-insoluble component added in subsequent steps. In a case where the ethanolinsoluble component added to the particles for supports in subsequent steps is confirmed in the separated ethanolinsoluble component, the average particle size of the particles for supports is obtained by eliminating the factors influencing the particle size distribution by the subsequent steps of addition. Specifically, the separation operation of the solvent-insoluble component is carried out by a properly selected solvent, or a combination thereof, so that the shape of the particles for supports can be confirmed after removing the surfactant composition and the components added in the subsequent steps.

[0174] The preferable properties of the detergent particles according to the present invention are as follows.
[0175] The bulk density is preferably from 500 to 1000 g/L, more preferably from 600 to 1000 g/L, especially preferably from 550 to 850 g/L.

[0176] The average particle size is preferably from 150 to 500 μm, more preferably from 180 to 400 μm.

11. Process for Preparing Detergent Particles

[0177] A preferable process for preparing detergent particles comprises the following step (I), and it may further comprise step (II) as occasion demands.

[0178] Step (I): mixing a surfactant composition with the particles for supporting a surfactant obtained in the process of the present invention, under condition that the surfactant composition is in a liquid or pasty state.

[0179] Step (II): mixing the mixture obtained in step (I) with a surface coating agent, thereby coating the surface of the powder detergent particles with the surface coating agent, provided that there is also included a case where step (II) proceeds simultaneously with the disintegration.

<Step (I)>

50

[0180] A process for supporting a surfactant composition by the particles for supports includes, for instance, a process comprising mixing the particles for supports with a surfactant composition by using a mixer for a batch process or continuous process. In the case of mixing by a batch process, as a process of supplying to a mixer, there may be employed such processes as (1) a process comprising previously supplying particles for supports in a mixer, and thereafter adding thereto a surfactant composition (;2) a process comprising uponlying particles for supports and a

surfactant composition in the mixer in small amounts at a time; (3) a process comprising supplying a part of particles for supports in a mixer, and thereafter supplying the remaining particles for supports and a surfactant composition in the mixer in small amounts at a time, and the like.

[0181] Among the surfactant compositions, those which are present as solids or pasty states even if heated within a practical temperature range, for instance, from 50° to 90°C, are previously dispersed or dissolved in a nonlonic surfactant having low viscosity, an aqueous solution of a nonlonic surfactant, or water, to prepare a liquid mixture or aqueous solution of a surfactant composition, to be added to the particles for supports in the form of a liquid mixture or aqueous solution. By this process, those surfactant compositions which are present as solids or pasty form can be easily added to the particles for supports. The mixing ratio of the surfactant composition having a low viscosity over to the solid or pasty surfactant composition is preferably such that the resulting liquid mixture or aqueous solution has a viscosity range of which is sorrayable.

[0182] The process for preparing the above liquid mixture includes, for instance, a process for mixing by supplying a solid or pasty surfactant composition to a surfactant having a low viscosity or water, or a process for preparing a liquid mixture of a surfactant composition by neutralizing an acid precursor of a surfactant, for instance, an acid precursor of an anionic surfactant, with an alkalizing agent, for instance, an aqueous sodium hydroxide or an aqueous potassium hydroxide, in a surfactant having a low viscosity or water.

[0183] Also, in this step, an acid precursor of an anionic surfactant can be also added before adding a surfactant composition, simultaneously with adding a surfactant composition, in the course of adding a surfactant composition or after adding a surfactant composition. By adding the acid precursor of an anionic surfactant, here can be aciding the acid precursor of an anionic surfactant, here can be aciding the acid precursor of an anionic surfactant, here can be aciding the acid provements in properties and quality, such as high concentration of the surfactants, supporting ability of particles for supports, control for the supporting ability thereof, and suppression of bleed-out of the nonionic surfactant and the flowability of the resulting determent naticles.

[0184] The acid precursor of an anionic surfactant which can be used in the present invention includes, for instance, alkylbenzeresulfonic acids, alkyle ther or alkenyl ether suffurio acids, alkyl- or alkyenylsulfurio acids, a-colefinsulfonic acids, or acids, and the like, it is especially preferable that the fatry acid is added after adding the surfactant, from the viewpoint of improvement in the flowability of the detergent particles.

[0185] The amount of the acid precursor of an anionic surfactant used is preferably from 0.5 to 30 parts by weight, more preferably from 1 to 20 parts by weight, is discovered by from 1 to 10 parts by weight, septically perfectly from 1 to 10 parts by weight, based on 100 parts by weight of the particles for supports. Here, the amount of the acid precursor used is not counted as the amount of the surfactant composition in the present invention. In addition, as the process for adding the acid precursor of an anionic surfactant, it is preferable that those in a liquid state at an ordinary temperature are supplied by spraying, and that those in a solid state at an ordinary temperature may be added as a powder, or they may be supplied by spraying after melting the solid. Here, in a case of adding the acid precursors as powder, it is preferable that the temperature of the detergent particles in the mixer is raised to a temperature at which the powder melts.

[0186] Preferable mixers are concretely as follows. In a case of mixing by a batch process, those of (1) to (3) are preferable: (1) Henschel Mixer (manufactured by Mitsul Milke Machinery Co., Ltd.); High-Speed Mixer (Fukae Powtec Corp.); Vertical Granulator (manufactured by Powrex Corp.); Lödige Mixer (manufactured by Matsuzaka Giken Co., Ltd.); PLOUGH SHARE Mixer (manufactured by PACIFIC MACHINERY & ENGINEERING Co., LTD.); mixers disclosed in Japanese Patent Laid-Open Nos. Hei 10-296064 and Hei 10-296065, and the like; (2) Ribbon Mixer (manufactured by Nichiwa Kikai Kogyo K.K.); Batch Kneader (manufactured by Satake Kagaku Kikai Kogyo K.K.); Ribocone (manufactured by K.K. Okawahara Seisakusho), and the like; (3) Nauta Mixer (manufactured by Hosokawa Micron Corp.). SV Mixer (Shinko Pantec Co., Ltd.), and the like. Among the above-mentioned mixers, preferable are Lödige Mixer. PLOUGH SHARE Mixer, and the mixers disclosed in Japanese Patent Laid-Open Nos. Hei 10-296064 and Hei 10-296065, and the like. Since step (II) described below can be carried out by the same mixer, these mixers are preferable from the viewpoint of simplification of equipments, Especially, the mixers disclosed in Japanese Patent Laid-Open Nos. Hei 10-296064 and Hei 10-296065 are preferable, because the moisture and temperature of the mixture can be regulated by agration, whereby the disintegration of the particles for supporting a surfactant can be suppressed In addition, mixers, such as Nauta Mixer, SV Mixer and Ribbon Mixer, which are capable of mixing powders with liquids without applying a strong shearing force, are preferable from the viewpoint that the disintegration of the particles for supporting a surfactant can be suppressed.

[0187] Also, the particles for supports may be mixed with a surfactant composition by using the above-mentioned continuous process-type mixer. Also, the continuous process-type mixer other than those listed above includes Flexo Mix (manufactured by Powery Corp.), Turbulizer (manufactured by Hosokawa Micron Corporation), and the like.

[0188] In addition, in this step, when a nonionic surfactant is used, it is preferable that a water-soluble nonionic organic compound (hereinafter referred to as "melting point-elevating agent") having a melting point of from 45° to 010°C and a molecular weight of from 1000 to 30000, or an aqueous solution thereof, which has a function of elevating

a melling point of this nonionic surfactant, can be added before adding a surfactant composition, simultaneously with adding a surfactant composition, in the course of adding a surfactant composition, or after adding a surfactant composition, or previously mixed with a surfactant composition. By adding the meting point-elevating agent, the caking property of the detergent particles and the bleed-out property of the surfactants in the detergent particles can be suppressed. Here, the same ones as those examplified in the melting point-elevating agent in the composition of the detergent particles described above can be used. The amount of the melting point-elevating agent used is preferably from 0.5 to 5 parts by weight, more preferably from 0.5 to 5 parts by weight, more preferably from 0.5 to 5 parts by weight, more preferably from 0.5 to 5 parts by weight, more preferably from 0.5 to 5 parts by weight, more preferably from 0.5 to 5 parts by weight, parts preferably from 1 to 3 parts by weight, based on 100 parts by weight of the particles for supports. The above range is preferable from the viewpoints of the suppression of the begregation between particles, the fast dissolubility, and the suppression of the bleed-out property and the caking property, each property of which is owned by the detergent particle contained in the detergent particles.

A process for adding the melting point-elevating agent, comprising adding by previously mixing the melting point-elevating agent, by a problemy process, or a process comprising adding a surfactant, and thereafter adding the melting point-elevating agent, is advantageous for the suppression of the bleed-out property and the caking property of the detergent particles.

- 5 (0189) As to the temperature within the mixer in this step, it is more preferable that mixing is carried out by healing to a temperature equal to or higher than the pour point of the surfactant, indentally, the pour point of the surfactant composition is measured according to the method of JIS K 2269. Here, the temperature to be heated may be a temperature higher than the pour point of the surfactant added in order to promote the support of the surfactant or position, and the practical temperature range is preferably from a temperature exceeding a pour point to 15 or 30°C. In addition, in the case where an acid precursor of an anionic surfactant is added in this step, it is more preferable to mix the components after heating to be imperature in which the acid precursor of an anionic surfactant can react.
- [0190] The mixing time in a batch process and the average residence time in the mixing in a confinuous process for obtaining the suitable detergent particles are preferably from 1 to 20 minutes, more preferably from 2 to 10 minutes. [0191] In addition, in the case where an equeous solution of a surfactant or an equeous solution of a water-soluble nonionic organic compound is added, a step of drying excess water contents during mixing and/or after mixing may be included.
- [0192] A powdery surfactant and/or a powdery builder can also be added before adding a surfactant composition, and the course of adding a surfactant composition, or after adding a surfactant composition. By adding the powdery builder, the particle size of the detergent particles can be controlled, and an improvement in detergency can be a chalved. Especially in the case where the acid precursor of an anionic surfactant is added, it is effective to add a powdery builder showing alkaline property prior to adding the acid precursor, from the viewpoint of accelerating the neutralization reaction. Incidentially, the term 'powdery builder' mentioned herein refers to an agent for enhancing detergency other than surfactants which is in a powdery form, concretely, including base materials showing metal ion capturing ability, such as zoolite and citrates; base materials showing capturing ability and alkikalizing ability, such as sodium carbonate; bese materials enhancing ionic strength, such as sodium suffact; and the like.
- [0193] Here, as crystalline silicates, those described in Japanese Patent Laid-Open No. Hei 5-278013, column 50, line 17 (sepecially, those prepared by a process comprising calicating and crystaltizing at a temperature of from 500° to 1000°C being preferable); Japanese Patent Laid-Open No. Hei 7-89712, column 2, line 45; and Japanese Patent Laid-Open No. Sho 60-27805, page 2, lower right column. Inter 16 (sepscially the silicates in Table 2 being preferable) can be used as powdery builders. Here, the alkali metal silicates having an SIO₂M₂O ratio, wherein M is an alkali metal, of from 0.5 to 32, preferably from 1.5 to 2.6 are favorably used.
- 5 [0194] The amount of the powdery builder used is preferably from 0.5 to 12 parts by weight, more preferably from 1 to 6 parts by weight, based on 100 parts by weight of the particles for supports. When the amount of the powdery builder for detergents used is in the above range, those having an excellent fast dissolubility are obtained.
 - [0195] Further, subsequent to step (I), it is preferable to add step (II) comprising surface-modifying the detergent particles.

<Step (11)>

[0196] In the present invention, in order to modify the particle surface of the detergent particles by which the surfactant is supported in step (I), the embodiments for addition may include a process comprising one or more steps of step (II) comprising adding various surface coeffing agents such as (1) fine powder, and (2) a liquid material.

[0197] Since the flowability and the anti-caking property of the detergent particles tend to improve by coating the particle surface of the detergent particles of the present invention, it is preferable to include a surface-modifying step. The devices used in step (II) are preferably those equipped with both aditation blades and disintegration blades among

the mixers exemplified in step (I). Each of the surface coating agents will be explained below.

(1) Fine Powder

[0198] As the fine powder, it is preferable that the average particle size of its primary particle is 10 µm or less, more preferably from 0.1 to 10 µm. When the particle size is in the above range, it is favorable from the viewpoints of the improvements in the coating ratio of the particle surface of the detergent particles, and improvements in the flowability and the anti-tasking property of the detergent particles. The average particle size of the fine powder can be measured by a method utilizing light scattering by, for instance; a particle analyzer (manufactured by Horbot, ITD.), or it may be measured by a microscopic observation or the like. In addition, it is preferable that the fine powder has a high lon exchange capacity or a high addikalizing ability from the sepect of detergency.

[0199] The fine powder is desirably aluminosilicates, which may be crystalline or amorphous. Besides them, fine powders of sodium sulfate, calcium silicate, silicon dioxide, bentonite, taic, day, amorphous silica derivatives, crystalline silicates, and the like are preferable. In addition, there can be also similarly used a metal soap of which primary particles have a size of 0.1 to 10 µm, a powdery surfactant (for instance, allysisulfates, and the like), or a water-soluble organic sail. In addition, when the crystalline silicate is used, it is preferably used in admixture with fine powder other than te crystalline silicate for the purpose of preventing deterioration owing to aggregation of the crystalline silicates by moisture absorption and carbon dioxide absorption, and the like.

[0200] The amount of the fine powder used is preferably from 0.5 to 40 parts by weight, more preferably from 1 to 30 parts by weight, especially preferably from 2 to 20 parts by weight, based on 100 parts by weight of the determinant particles, When the amount of the fine powder used is in the above range, the flowebility is improved, thereby giving a good sense of feet to consumers.

(2) Liquid Materials

[0201] The liquid materials include water-soluble polymers, fatty acids, and the like, which may be added in the form of aqueous solutions and molten states.

(2-1) Water-Soluble Polymer

[0202] The water-soluble polymer includes carboxymethyl celluloses, polyethylene glycols, polycarboxylates such as sodium polyacrylates and copolymers of ecryl acid and maleic acid and salts thereof, and the like. The amount of the water-soluble polymer used is preferably from 0.5 to 10 parts by weight, nor preferably from 1 to 8 parts by weight, especially preferably from 2 to 6 parts by weight, based on 100 parts by weight of the detergent particles. When the amount of the water-soluble polymer used is in the above range, the detergent particles exhibiting excellent dissolubility and excellent flowability and enti-caking properties can be obtained.

(2-2) Fatty Acid

30

[0203] The fatty acid includes, for instance, fatty acids having 10 to 22 carbon atoms, and the like. The amount of the fatty acid used is preferably from 0.5 to 5 parts by weight, especially preferably from 0.5 to 3 parts by weight, based on 100 parts by weight of the detergent particles. In a case of a fatty acid in a solid state at ordinary temperature, it is preferable that the fatty acid is heated to a temperature exhibiting flowability, and then supplied to the detergent particles by spraying.

12. Detergent Composition

[0204] The detergent composition in the present invention is a composition comprising the detergent particles described above, and the composition further comprises separately added detergent components other than the detergent particles for instance, builder particles, fluorescent dyes, enzymes, perfumes, deforming agents, bleaching activators, and the like).

(0205) The content of the detergent particles in the detergent composition is preferably 50% by weight or more, more preferably 80% by weight or more, still more preferably 70% by weight or more, still more preferably 80% by weight or more, especially preferably 70% by weight.

(0206) The content of the detergent components other than the detergent particles in the detergent composition is preferably 50% by weight or less, more preferably 40% by weight or less, still more preferably 30% by weight or less, especially preferably 20% by weight or less.

13. Method for Measurement of Properties

10207] The values for the properties in the present specification are measured by the following methods. (Bulk Density): measured by a method according to JIS K 3362. (Average Particle Size): measured using standard sieves according to JIS 2 8011. For example, nine-step sieves each having a sieve-opening of 2000 μm, 1400 μm, 1000 μm, 710 μm, 500 μm, 355 μm, 250 μm, 180 μm, and 125 μm, and a receiving tray are used, and the sieves and the receiving tray are attached to a rotating and tapping shaker machine (mandratured by HEIKO SEISANLSHO), tapping: 156 times/min, rolling: 290 times/min). A 100 g sample is vibrated for 10 minutes to be classified. Thereafter, the mass base frequency is sequentially cumulated for each of sieve-on particles in the order of the receiving tray, and sieve-brening of a first sieve of which cumulative mass base frequency is 50% or more is defined as μ μm, and a sieve-opening of one sieve-opening agree than a μm is defined as β μm, in the case where the cumulative mass base frequency from the receiving tray to the α μm-sieve is defined as β, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the mass base frequency from the receiving tray to the α μm-sieve is defined as 6 km, and the receiving tray to the α μm-sieve is defined as 6 km, a

(Average particle size) = 10^A:

wherein

15

20

25

40

$$A = \frac{(\gamma - \frac{\theta}{\log \beta - \log \alpha} \times \log \beta)}{\frac{\theta}{\log \beta - \log \alpha}}$$

(Particle Strength): A cylindrical vessel of an inner diameter of 3 cm and a height of 8 cm is charged with 20 g of a sample, and the sample-containing vessel (manufactured by Tsutsui Rikagaku Kikai K.K., "Model TVP1" tapping-type close-packed bulk density measurement device; tapping conditions: period 36 times/minute, free flow from a height of 60 mm) is tapped for 30 times. The sample height (an initial sample height) at that time is measured. Thereafter, an entire upper surface of the sample kept in the vessel is pressed at a rate of 10 mm/min with a pressing machine to take measurements for a load-displacement curve. The slope of the linear portion at a displacement rate of 5% or less is multiplied by an initial sample height, and the resulting product is divided by a pressed area, to give a quotient which is defined as particle strength. (Supporting Capacity and Supporting Rate of Liquid Surfactant Composition): A cylindrical mixing vessel of an inner diameter of 5 cm and a height of 15 cm which is equipped with aditation impellers in the inner portion thereof is charged with 100 g of the particles. With stirring the contents at 350 rpm, a polyoxyethylene alkyl ether (C12/C14 = 6/4; EO = 7.7; melting point: 25°C) is added dropwise at 30°C at a rate of 10 mL/min, and the change of agitation torque with the passage of time is measured. A value obtained by dividing the amount of the polyoxyethylene alkyl ether supplied at a point where the agitation torque reaches the highest level by the weight (100 g) of the particles is defined as the supporting capacity (mL/g) of the particles. In addition, the supporting rate is expressed as a value (mL/g) obtained by dividing the amount of the polyoxyethylene alkyl ether supplied at a point where the amount of change per unit time is the largest in the process of increasing the agitation torque until the agitation torque exhibits the highest level by the weight (100 g) of the particles. The larger this numerical value, the more excellent the supporting rate, i.e. the more excellent the supporting rate of the particles, the more suppressed the excess polyoxyethylene alkyl ether is suppressed, thereby resulting in a delay in the time period for which the agitation torque is increased. (Water Content): The water content of the particles is measured by infrared moisture meter method. Specifically, a 3 g sample is weighed and placed on a weighing dish of a known weight, and the sample is heated and dried for 3 minutes with an infrared moisture meter (manufactured by Kett Kagaku Kenkyujo K.K. (infrared rav lamp: 185 W)). After drying, the dried sample and the weighing dish are weighed. The water content in the sample is calculated by taking the difference in the weights of the container and the sample before and after drying obtained by the above operation, dividing the difference with the weight of the sample weighed, and multiplying the results with 100. (Microporous Capacity Distribution): The microporous capacity of the particles for supporting a surfactant is determined as follows by using mercury porosimeter, "manufactured by SHIMADZU CORPORATION, "Poresizer 9320") in accordance with its instruction manual. Specifically, a cell is charged with 200 mg of particles for supporting a surfactant and the pressed mercury is measured separately for a low-pressure portion (0 to 14.2 psia) and a high-pressure portion (14.2 to 30000 psia). The leveling of the measurement data is carried out by taking an average of two each to obtain a mode diameter of 0.01 to 3 µm and a microporous capacity.

(Flowability): The flow time refers to a time period required for flowing 100 mL of detergent powder from a hopper used in a measurement of bulk density as defined in JIS K 3362.

(Anti-Caking Property): An open-top carton having dimensions of 10 cm in length, 6 cm in width, and 4 cm in height is made out of a Model No. 2 filter paper defined by JIS P 3801 (for instance, qualitative No. 2 filter paper, manufactured by Toyo Roshi K.K.). A 100 g sample is placed in this carton, and an acrylic resin plate and a lead plate (or an iron plate) with a total weight of 15 c + 250 o are placed on the sample. The above carton is maintained in a thermostat

5 kept at a temperature of 30°C and at a humidity of 80%, and the caking conditions after 7 days are evaluated as explained below. The evaluation is made by obtaining the permeability as follows. The higher the permeability, the higher the anti-caking property, which is preferable properties as the detergent particles.

(Permeability): A sample obtained after the above test is gently placed on a sieve (sieve opening: 4760 µm, as defined by JIS Z 8801), and the weight of the powder passing through the sieve is measured. The permeability based on the sample after the test is calculated.

(Bleed-Out Property): An open-top carton having dimensions of 10 cm in length, 6 cm in width, and 4 cm in height is made out of a Model No. 2 filter paper defined by JIS P 3801 (for instance, qualitative No. 2 filter paper, manufactured by Toyo Roshi K.K.). A line with a width of 0.5 to 1.0 mm is diagonally drawn on the bottom surface of the carton, which is the surface of the packed sample, using a Magic Marker (manufactured by K.K. UCHIDA YOKO, "Magic Ink M700-T1"). A 100 g sample is packed in this carton, and an acrylic resin plate and a lead plate (or an iron plate) with a total weight of 15 g + 250 g are placed on the sample. The carton is placed in a moisture-proof carton, and allowed to stand in a thermostal kept at a temperature of 30°C. After 7 days, the bleed-out property was evaluated by visually examining the decree of blur of the Magic Marker. The evaluation criteria are as follows.

20 Rank 5: Blurred width of the Magic Marker being 2 cm or more.

Rank 4: Blurred width of the Magic Marker being 1 cm or more.

Rank 3: Blurred width of the Magic Marker being 0.5 cm or more.

Rank 2: Slight blur of the Magic Marker being found.

Rank 1: No blur of the Magic Marker being found.

14. Process for Preparing Detergent Composition

[0208] The process for preparing a detergent composition is not particularly limited, and an example thereof include a process of mixing the detergent particles and separately added detergent components. Since the detergent composition obtained in the manner described above contain a detergent particle having a large supporting capacity the surfactant, sufficient detergent effects can be exhibited even with a small amount. The application of such a detergent composition is not particularly limited, as long as it is applied to powder detergent, including, for instance, laundry powder detergents, detergents for dishwasher, and the like.

35 EXAMPLES

[0209] In the present examples, the following starting materials were used unless otherwise specified. Sodium sulfate: anhydrous neutral sodium sulfate (manufactured by Shikoku Kasei K.K.)

Sodium sulfite: sodium sulfite (manufactured by MITSUI CHEMICALS, INC.)

Fluorescent dye: Tinopal CBS-X (manufactured by Ciba Specialty Chemicals)

Sodium carbonate: DENSE ASH (average particle size: 290 µm; manufactured by Central Glass Co., Ltd.)

40% By weight aqueous solution of sodium polyacrylate: weight-average molecular weight: 10000 (manufactured by Kao Corporation)

Sodium chloride: roast salt S (manufactured by Nippon Seien K.K.)

45 Crystalline sodium aluminosilicate (zeolite): TOYOBUILDER (4A type; average particle size: 3.5 μm) (manufactured by Tosoh Corporation) Potyoxyethylene alkyl ether: EMULGEN 108 KM (average moles of ethylene oxides: 8.5; number of carbon atoms in alkyl molety: 12 to 14; manufactured by Kao Corporation.

Polyethylene glycol: K-PEG 6000 (weight-average molecular weight: 8500; manufactured by Kao Corporation)

Amorphous aluminosilicate: a product prepared by pulverizing the composition of Preparation Example 2 described in Japanese Patent Laid-Open No. Hel 9-132794 to an average particle size of 8 um.

Example 1

[0210] A mixing vessel was charged with 375 parts by weight of water. After the water temperature reached 35°C, 127 parts by weight of sodium sulfate, 5 parts by weight of sodium sulfite, and 1 part by weight of a fluorescent dye were added thereto, and the resulting mixture was agitated for 10 minutes. One-hundred and twenty-seven parts by weight of sodium carbonate were added to the mixture, and 75 parts by weight of a 40% by weight acqueous solution of sodium polyacytale were added thereto. The resulting mixture was agitated for 10 minutes, to give a first preparation.

liquid. Twenty-four parts by weight of sodium chloride, a microcrystal-precipitating agent, were added thereto, and the resulting mixture was agitated for 10 minutes. Further, 266 parts by weight of zeolite were added, and the resulting mixture was agitated for 30 minutes, to give a homogenous second preparation liquid (water content of slurry: 42%) by weight). The final temperature of this preparation liquid was 40°C. The amount of the water-soluble inorganic sait precipitated by the addition of sodium chloride was 16.3% by weight) of that dissolved in the first preparation liquid [0211] After the preparation of the first preparation liquid and 10 minutes after the addition of sodium chloride, a

[0211] After the preparation of the first preparation liquid and 10 minutes after the addition of sodium chloride, a sample was taken from each of the preparation liquids, and the number of particles and the particle size distribution were determined by TSUB-TEC M100.

[0212] The number of particles in the first preparation liquid was 778 counts/s, and the average particle size (on a number basis) was 172 µm. The number of particles in the second preparation liquid after the addition of sodium chloride was 2634 counts/s, and the average particle size was 21.2 µm. From these determination results, the number of water-soluble salt was increased by 1856 counts/s by the addition of sodium chloride, and the average particle size of the increased water-soluble salt was 1.6 µm.

[0213] The second preparation liquid was fed to a spray-drying tower (countercurrent flow type) by a pump, and sprayed from a pressure-spray nozzle attached near the top of the tower at a spraying-pressure of 2.5 MPa. The high-temperature gas to be fed to the spray-drying tower was fed at a temperature of 200°C from the bottom of the tower, and exhausted at 80°C from the top of the tower. The water content of the resulting Particles for Supporting Surfact 1 was 4% by weight. Detergent Particles 1 were prepared using Particles for Supporting Surfactant 1 by the method show below.

[0214] A surfactant composition (polyoxyethylene alkyl ether/polyethylene glycol/sodium alkylbenzenesulfonate/water = 42/84/216 (weight ratio)) was adjusted to 80°C. Next, 100 parts by weight of the resulting Particles for Supporting Surfactant 1 were supplied into a Lödige Mixer (manufactured by Matsuzaka Géhen Co., Lid.; espacity; 130 L; equipped with a jacket), and the agitation of a main shaft (agitation impeliers: rotational speed: 60 pm; peripheral speed: 1.6 m/ s) was started. Incidentally, hot water at 80°C was allowed to flow through the jacket at 10 Liminute. Fifty parts by weight of the above surfactant composition were supplied into the above mixer in 2 minutes, and thereafter the resulting mixture was agitated for 5 minutes. Further, 6 parts by weight, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, of an amorphous aluminosilicate were supplied thereinto. The agi-

tations of the main shaft (rotational speed: 120 rpm; peripheral speed: 3.1 m/s) and a chopper (rotational speed: 3600 rpm; peripheral speed: 28 m/s) were carried out for 1 minute, and Detergent Particles 1 were discharged.

Example 2

20

[0215] Particles for Supporing Surfactant 2 were obtained in the same manner as in Example 1, Detergent Particles 2 were prepared in the same manner as in Example 1 using Particles for Supporting Surfactant 2. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 4 parts by weight.

Comparative Example 1

70 [0216] Particles for Supporing Surfactant 3 were obtained in the same manner as in Example 1 except that sodium chloride, a microcrystal-precipitating agent, was added prior to the addition of a water-soluble salt, and agitated for 10 minutes to be completely dissolved. Detergent Particles 3 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 3. However, in the case where the amorphous aluminosilicate was used in an amount of 6 parts by weight, the same amount as that of Example 1, Particles for Supporting Surfactant 3 of oil not sufficiently support the surfactant composition and became aggregated during the agitation in a Lödige Mixer, so that the values of the properties were deteriorated to an extent to be undeterminable.

Example 3

10217] Particles for Supporting Surfactant 4 were obtained in the same manner as in Example 1 except that sodium bromide (manufactured by OTSURA CHEMICAL CO., LTD) was used as a microcrystal-precipitating agent. The amount of the water-soluble incorpain call precipitating of sodium bromide was 2.7% by weight of that dissolved in the first preparation liquid. Detergent Particles 4 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 4. The amount of an amount of surfacet as 1, usar 7 parts by weight amount in which the bleed-out property the detergent particles is to be evaluated as 1, usar 7 parts by weight.

Comparative Example 2

30

45

50

55

[0218] Particles for Supporting Surfactant 5 were obtained in the same manner as in Comparative Example 1 except hat sodium bromide (manufactured by OTSUKA CHEMICAL CO., LTD) was used as a microcrystal-precipitating agent. Detergent Particles 5 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 5. However, in the case where the amorphous aluminosilicate was used in an amount of 7 parts by weight, the same amount as that of Example 3. Particles for Supporting Surfactant 5 din ont sufficiently support the surfact composition and became aggregated during the agitation in a Lödige Mixer, so that the values of the properties were deteriorated to an extent to be undeterminated to

[0219] The composition and the properties of each group of the resulting Particles for Supporting Surfactant 1 to 5 are shown in Table 1, and the proparties of each group of Detergent Particles 1 to 6 are shown in Table 2. In Examples of the present invention, the particle size of the water-soluble salt precipitated in the sturry is made fine due to the effect of the microcystal-precipitating agent, in addition, by increasing the amount of the microcystal-precipitating agent, more water-soluble salt can be precipitated. Therefore, the particles for supporting a surfactant of the precipitating agent, more water-soluble salt can be precipitated. Therefore, the particles for supporting a surfactant of the precipitation in the supporting Surfactant 1, 2 and 4) have a smaller mode diameter of microporous capacity distribution than that in Comparative Examples, thereby having a microporous capacity distribution attracting comparison of the present invention (each group of Detergent Particles 1, 2 and 4), the amount of the amorphous aluminositicate could be reduced.

Tohlo 1

Table 1								
	Ex. 1	Ex. 2	Comp. Ex. 1	Ex. 3	Comp. Ex. 2			
Composition % By Weight								
Zeolite	44.0	42.0	44.0	44.0	44.0			
Sodium Carbonate	21.0	20.0	21.0	21.0	21.0			
Sodium Sulfate	21.0	20.0	21.0	21.0	21.0			
Sodium Sulfite	0.8	0.8	0.8	0.8	0.8			
Sodium Polyacrylate	5.0	5.0	5.0	5.0	5.0			
Fluorescent Dye	0.2	0.2	0.2	0.2	0.2			
Sodium Chloride	4.0	8.0	4.0	0.0	0.0			
Sodium Bromide	0.0	0.0	0.0	4.0	4.0			
Water	4.0	4.0	4.0	4.0	4.0			
TOTAL .	100.0	100.0	100.0	100.0	100.0			
Operation	Ī .							
Post-Addition of Microcrystal-	0	0		0				
Precipitating Agent								
Concentration Operation								
Precipitation by Temperature								
Adjustment	l							
Slurry Pulverization								
Slurry								
Water Content of Slurry [%]	42	42	42	42	42			
Temperature of Slurry [°C]	40	40	40	40	40			
Increased Amount of	16.3	32.5		3.0	-			
Undissolved Salt [%]								
Particle Properties								
Average Particle Size [µm]	250	253	245	240	242			
Bulk Density [g/L]	601	603	599	607	610			
Particle Strength [MPa]	29	34	28	32	30			
Supporting Capacity [mL/g]	0.45	0.51	0.44	0.38	0.37			
Mode Diameter of Microporous	0.81	0.67	1.63	0.78	1.58			
Capacity Distribution [µm]								

Table 1 (continued)

	Ex. 1	Ex. 2	Comp. Ex. 1	Ex. 3	Comp. Ex. 2	Ì
Particle Properties						l
0.01 - 3 μm [mL/g]	0.32	0.34	0.29	0.33	0.29	ĺ

Table 2

Table 2								
	Ex. 1	Ex. 2	Comp. Ex. 1	Ex. 3	Comp. Ex. 2			
Composition of Detergent Parts by Weight								
Particles for Supporting Surfactant 1	100							
Particles for Supporting Surfactant 2	l	100		l				
Particles for Supporting Surfactant 3		l	100					
Particles for Supporting Surfactant 4				100				
Particles for Supporting Surfactant 5		İ			100			
Surfactants (Sodium Alkylbenzenesulfonate)	50	50	50	50	50			
	(21)	(21)	(21)	(21)	(21)			
(Polyoxyethylene Alkyl Ether)	(21)	(21)	(21)	(21)	(21)			
(Polyethylene Glycol)	(4)	(4)	(4)	(4)	(4)			
(Water)	(4)	(4)	(4)	(4)	(4)			
Amorphous Aluminosilicate	6	4	6	7	11			
Propertiés								
Average Particle Size [µm]	258	264	Undeterminable	262	Undeterminabl			
Bulk Density [g/L]	738	748	Undeterminable	745	Undeterminabl			
Flowability [s]	6.2	6.1	Undeterminable	6.2	Undeterminabl			
Bleed-out Property	1	1	5	1	5			

Example 4

5

10

15

20

25

30

[0220] A mixing vessel equipped with a jacket comprising a pressure-reducing device and an agitator, was charged with \$15 parts by weight of water and the temperature was raised to 35°C. One-hundred and eight parts by weight of sodium suffile, \$6 parts by weight of sodium suffile, \$6 parts by weight of a 40% by weight acueous solution of sodium polyacryste, 1 part by weight of afluorescent dye, and 205 parts by weight of a fluorescent dye, and 205 parts by weight of a fluorescent dye, and 205 parts by weight of a fluorescent dye, and 205 parts by weight of a fluorescent dye, and 205 parts by weight of a fluorescent dye, and 205 parts by weight of a fluorescent dye, and 205 parts by weight of a fluorescent dye and the
[0221] Water was evaporated, with heating the first preparation liquid by allowing hot water at 65°C to flow through the jacket under a reduced pressure of 100 Torr, to concentrate the fiquid to a water content of 45% by weight. The amount of the water-soluble inorganic salt (average particle size: 18 µm) precipitated by the concentration operation was 25% by weight of that dissolved in the first preparation liquid.

[0222] The concentrated second preparation liquid was spray-dried in the same manner as in Example 1. The hightemperature gas to be supplied to the spray-drying tower was fed at a temperature of 220°C from the bottom of the tower, and exhausted at 110°C from the top of the tower. The water content of the resulting Particles for Supporting Surfactant 6 was 4% by weight.

[0223] Detergent Particles 6 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 6, The amount of an amorphous alluminosillicate supplied, as the minimum amount in which the bleed-out property the detergent particles is to be evaluated as 1, was 1.5 parts by weight.

Example 5

55

[0224] Particles for Supporting Surfactant 7 were obtained in the same manner as in Example 4 except that a first preparation liquid having a water content of 50% by weight was prepared by adjusting the amount of water to be added, and that a second preparation liquid was obtained by concentrating the first preparation liquid by a water content of the content of the second preparation liquid was obtained by concentrating the first preparation liquid by a water content.

45% by weight. The amount of the water-soluble inorganic salt (average particle size: 20 µm) precipitated in the second preparation liquid was 19% by weight of that dissolved in the first preparation liquid.

[0225] The number of particles and the particle size distribution before and after the concentration in the preparation liquid were determined by TSUB-TEC M100. Inciderallally, in order to increase the accuracy of the determination field the determination was carried out using a liquid (water content of slurry; 64.9% by weight) corresponding to a first preparation liquid prepared in a separate mixing vessel without blending zeolite, and a liquid (water content of slurry; 63.1% by weight) corresponding to a second preparation liquid prepared by concentrating the liquid corresponding to a stret preparation liquid. The number of particles in the liquid corresponding to a first preparation liquid. The number of particles in the liquid corresponding to a first preparation liquid was 426 counts/s, and the average particle size (on a number basis) was 114 µm. The number of particles in the liquid corresponding to a second preparation liquid after the concentration was 8351 counts/s, and the average particle size was 20.0 µm. From these determination results, the number of particles of the water-soluble sett was increased by 5925 counts/s.

the concentration, and the average particle size of the increased water-soluble salt was 18.5 µm. [0226] Detergent Particles 7 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 7. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 2.5 parts by weight.

Comparative Example 3

[0227] Particles for Supporting Surfactant 8 were obtained in the same manner as in Example 4 except that a preparation liquid having a water content of 45% by weight was prepared by adjusting the amount of water to be acided, and that the concentration was not carried out. Detergent Particles 8 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 8. The amount of an amorphous aluminosilicate supplied, as the mininum amount in which the bleed-out property of the detergent particles to be evaluated as 1, was 8 paired, as the mininum amount of less than 8 parts by weight. In the case where the amorphous aluminosilicate was used in an amount of less than 8 parts by weight, the bleed-out property evaluated as 1 was not obtained.

Comparative Example 4

[0228] Particles for Supporting Surfactant 9 were obtained in the same manner as in Example 4 except that a preparation liquid having a water content of 55% by weight was prepared by adjusting the amount of water to be added, and that the concentration was not carried out. The water-soluble components in the preparation liquid were completely dissolved. Detergent Particles 9 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 9. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 6 parts by weight. In the case where the amorphous aluminosilicate was used in an amount of less than 6 parts by weight, the bleed-out property evaluated as 1 was not obtained.

Example 6

[0229] A first preparation liquid was prepared in the same manner as in Example 4, and concentrated to a water content of 46% by weight. Subsequently, 19 parts by weight of sodium chloride, a microcrystal-precipitating agent, were further added thereto, and thereafter the resulting mixture was agitated for 30 minutes, to give a second preparation liquid (water content: 45% by weight). The amount of the water-sobble inorganic salt precipitated by the concentration operation and the addition of the microcrystal-precipitating agent was 35.7% by weight of that dissolved in the first orderaration floud;

[0230] The second preparation liquid was spray-dried in the same manner as in Example 1, to give Particles for Supporting Surfactant 10.

[0231] Detergent Particles 10 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 10. Detergent Particles 10 had a sufficiently excellent flowability, and the level of the bleed-out property was evaluated as 1 without addition of an amorphous aluminositicate.

Example 7

(0232) A first preparation liquid was prepared in the same manner as in Example 6, and thereafter Particles for Supporting Surfactant 11 were obtained in the same manner as in Example 6. Detergent Particles 11 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 11. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 1 part by weight.

[0233] The composition and the properties of each group of the resulting Particles for Supporting Surfactant 8 to 11 are shown in Table 4. The properties of each group of Detergent Particles 8 to 11 are shown in Table 4. [0234]. From the results shown in Table 4 and 4, since each group of Particles for Supporting Surfactant 8 and 9 has a relatively time are many forms.

has a relatively low supporting ability, it was necessary to add a large amount of an amorphous aluminosilicate when trying to obtain detergent particles having an excellent bleed-out properly using the particles. [0236] On the other hand, since each group of Particles for Supporting Surfactant 8 and 7 obtained by the concen-

tration operation has a mode diameter of microporous capacity distribution of 1.5 µm or less and a high supporting ability, detergent particles having an excellent bleed-out properly could be obtained by using these groups of the particles, even in the case where the amount of the amorphous aluminositiate was reduced. In addition, the supporting ability of the particles for supporting a surfactant could be further improved by carrying out both a concentration operation of surry and addition of a microcrystal-precipitating agent.

Table 3

15		Ex. 4	Ex. 5	Comp. Ex. 3	Comp. Ex. 4	Ex. 6	Ex. 7
	Composition % By Weight					1	
	Zeolite	44.0	44.0	44.0	44.0	40.0	40.0
	Sodium Carbonate	23.0	23.0	23.0	23.0	23.0	23.0
	Sodium Sulfate	23.0	23.0	23.0	23.0	23.0	23.0
20	Sodium Sulfite	0.8	0.8	0.8	0.8	0.8	0.8
	Sodium Polyacrylate	5.0	5.0	5.0	5.0	5.0	5.0
	Fluorescent Dye	0.2	0.2	0.2	0.2	0.2	0.2
	Sodium Chloride	0.0	0.0	0.0	0.0	4.0	4.0
25	Sodium Bromide	0.0	0.0	0.0	0.0	0.0	0.0
	Water	4.0	4.0	4.0	4.0	4.0	4.0
	TOTAL	100.0	100.0	100.0	100.0	100.0	100.0
	Operation						
30	Post-Addition of Microcrystal-					0	0
	Precipitating Agent						
	Concentration Operation	0	0			0	0
	Precipitation by Temperature						
25	Adjustment						
	Slurry Pulvenzation						
	Slurry						
	Water Content of Slurry [%]	55>45	50-→45	45	55	5545	50→45
o	Temperature of Slurry [°C]	60	60	60	60	60	60
	Increased Amount of Undissolved Salt [%]	25	19	-	-	35.7	27
	Particle Properties			1			
5	Average Particle Size [µm]	280	265	235	210	264	258
-	Bulk Density [g/L]	615	600	600	480	601	605
	Particle Strength [MPa]	28	28`	28	17	30	30
	Supporting Capacity [mL/g]	0.68	0.6	0.42	0.53	0.66	0.64
	Mode Diameter of Microporous	0.82	0.96	1.8	2.2	0.55	0.54
0	Capacity Distribution [μm] 0.01 - 3 μm [mL/g]	0.37	0.36	0.35	0.47	0.38	0.37

Table 4

		DIC 4				
	Ex. 4	Ex. 5	Comp. Ex. 3	Comp. Ex. 4	Ex. 6	Ex. 7
Composition of Detergent Parts by Weight	1					
Particles for Supporting Surfactant 6	100					
Particles for Supporting Surfactant 7	1	100				
Particles for Supporting Surfactant 8	l		100			
Particles for Supporting Surfactant 9	l			100		
Particles for Supporting Surfactant 10	l				100	
Particles for Supporting Surfactant 11						100
Surfactants	50	50	50	50	50	50
(Sodium Alkylbenzenesulfonate)	(21)	(21)	(21)	(21)	(21)	(21)
(Polyoxyethylene Alkyl Ether)	(21)	(21)	(21)	(21)	(21)	(21)
(Polyethylene Glycol)	(4)	(4)	(4)	(4)	(4)	(4)
(Water)	(4)	(4)	(4)	(4)	(4)	(4)
Amorphous						
Aluminosilicate	1.5	2.5	8	6	0	1
Properties	ļ					
Average Particle Size [µm]	300	280	245	230	271	268
Bulk Density [g/L]	740	740	730	660	742	743
Flowability [s]	6.3	6.3	6.3	6.4	6.3	6.3
Bleed-out Property	1	1	1	1	1	1

Example 8

5

10

15

20

35

50

[0236] A mixing wessel equipped with a jacket, comprising an agilator, was charged with 407 parts by weight of water, and hot water at 40°C was allowed to flow through the jacket. One-hundred and thirty-two parts by weight of sodium suffie, and 1 part by weight of a fluorescent dye were added thereto, and the resulting mixture was agitated for 10 minutes. One-hundred and thirty-two parts by weight of sodium carbonate were added to the mixture, and 2 parts by weight of a 40% by weight acquous solition of sodium polyacrylate and 252 parts by weight of zoolite were sequentially added thereto. The resulting mixture was agitated for 15 minutes, to give a first preparation liquid at 40°C and 100°C and 1

[0237] Next, hot water at 60°C was allowed to flow through the jocket, and the liquid mixture was agitated for 30 minutes, thereby adjusting the temperature of the preparation liquid to 60°C, to give a second preparation liquid. Then viscosity of the preparation liquid was increased from 60 mPs = to 1200 mPs = 50 the heating operation. The amount of the water-soluble inorganic salt precipitated by the operation was 8.2% by weight of that dissolved in the first preparation flow.

[0238] The resulting second preparation liquid was spray-dried in the same manner as in Example 1. The hightemperature gas to be fed to the spray-drying lower was fed at a temperature 0.210°C from the bottom of the tower, and exhausted at 105°C from the top of the tower. The water content of the resulting Particles for Supporting Surfactant 12 was 4% by weight.

[0239] Detergent Particles 12 were prepared in the same manner as in Example 1 using the resulting Particles for Sopporting Surfactant 12. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 6 parts by weight.

Example 9

[0240] A first preparation liquid at 40°C was prepared under the same procedures as in Example 8. The preparation liquid was allowed to flow through a shell and tube-type heat exchanger, thereby raising the temperature of the preparation liquid. In the preparation liquid, not prejaration indirecrystates of microcrystates of microcryst

10241] The number of particles and the particle size distribution before and after the concentration in the preparation liquid were determined by TSUB-TEC M100. Incidentally, the determination was carried out in the same manner as in Example 4, using a liquid corresponding to a first preparation liquid (water content of sury; 6.01% by weight) prepared in a separate mixing vessel without blending zeolite, and a liquid corresponding to a second preparation liquid to 70°C. The number of particles in the liquid corresponding to a first preparation liquid vary 70°C. The unitaber of particles in the liquid corresponding to a second preparation liquid after raising the temperature was 25°C counts/s, and the average particle size was 26°C Jum. From these determination results, the number of particles of the water-soluble salt was increased by 746°C counts/s by the heating operation, and the average particle size of the increased water-soluble salt was increased by 746°C counts/s by the heating operation, and the average particle size of the increased water-soluble salt was increased by 746°C counts/s by the heating operation, and the average particle size of the increased water-soluble salt was increased by 746°C counts/s by the heating operation, and the average particle size of the increased water-soluble salt was increased water.

[0242] The resulting second preparation liquid was spray-dried in the same manner as in Example 1. The hightemperature gas to be fed to the spray-drying tower was fed at a temperature of 220°C from the bottom of the tower, and exhausted at 110°C from the top of the tower. The water content of the resulting Particles for Supporting Surfactant 2 was 4% by weight.

[0243] Detergent Particles 13 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 13. The amount of an amorphous aluminositicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 5 parts by weight.

Comparative Example 5

20

30

[0244] A first preparation liquid at 40°C was prepared under the same procedures as in Example 8, and the preparation liquid was spray-dried under the same conditions as in Example 8 without heating the preparation liquid, to give Particles for Supporting Surfactant 14. Detergent Particles 14 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 14. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles its to be evaluated as 1, was 8 parts by weight. When the amorphous aluminosilicate was used in an amount of less than 8 parts by weight, the bleed-out property evaluated as 1 was not obtained.

Comparative Example 6

[0245] Particles for Supporting Surfactant 15 were prepared in the same manner as in Comparative Example 5 except that a first preparation fiquid at 70°C was obtained by changing the temperature of hot water to be allowed to flow into the lacket to 70°C. Detergent Particles 15 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 15. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 10 parts by weight. When the amorphous aluminosilicate was used in an amount of less than 10 parts by weight, the bleed-out property evaluated as 1 was not obtained.

Example 10

[0246] A first preparation liquid was prepared in the same manner as in Example 9. Next, the slurry was allowed to flow into a shell and tube-type heat exchanger, thereby raising the temperature of the preparation liquid to 70°C. Thereafter, a microcrystal-precipitating agent was further added thereto, to give a second preparation liquid. The amount of the water-soluble inorganic salt precipitated by the heafing operation of the preparation liquid was 25.2% by weight of the amount dissolved in the first proparation liquid.

[0247] The resulting second preparation liquid was spray-dried in the same manner as in Example 1. The hightemperature gas to be fed to the spray-drying tower was fed at a temperature of 205°C from the bottom of the tower, and exhausted at 95°C from the top of the lower. The water content of the resulting Particles for Supporting Surfactant 16 was 4% by weight.

60 [0248] Delergent Particles 16 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 16. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is evaluated as 1, was 3 parts by weight.

[0248] The composition, the properties and the like of each group of the resulting Particles for Supporting Surfactant 12 to 16 are shown in Table 5, and the properties of each group of Detrepent Particles 12 to 16 are shown in Table 6, 10250] From the results shown in Tables 5 and 6, since each group of Particles for Supporting Surfactant 14 and 15 has a relatively low supporting ability, twa necessary to add a large amount of an amorphous aluminosilicate when trying to obtain detergent particles, having an excellent bleed-out property using the particles.

[0251] On the other hand, since each group of Particles for Supporting Surfactant 12 and 13 has a mode diameter

of microporous capacity distribution of 1.5 μ m or less and a high supporting ability, detergent particles having an excellent bleed-out property could be obtained by using these groups of the particles for supporting a surfactant, even when the amount of the amorphous aluminositicate was reduced.

[0252] In addition, the supporting ability of the particles for supporting a surfactant could be further improved by carrying out both a concentration operation of slurry and addition of a microcrystal-precipitating agent.

10

15

25

30

35

50

55

Ta		

	Tab	e 5			
	Ex. 8	Ex. 9	Comp. Ex. 5	Comp. Ex. 6	Ex. 10
Composition % By Weight					
Zeolite	44.0	44.0	44.0	44.0	40.0
Sodium Carbonate	23.0	23.0	23.0	23.0	23.0
Sodium Sulfate	23.0	23.0	23.0	23.0	23.0
Sodium Sulfite	0.8	0.8	0.8	0.8	8.0
Sodium Polyacrylate	5.0	5.0	5.0	5.0	5.0
Fluorescent Dye	0.2	0.2	0.2	0.2	0.2
Sodium Chloride	0.0	0.0	0.0	· 0.0	4.0
Sodium Bromide	0.0	0.0	0.0	0.0	0.0
Water -	4.0	4.0	4.0	4.0	4.0
TOTAL .	100.0	100.0	100.0	100.0	100.0
Operation					
Post-Addition of Microcrystal-	1				0
Precipitating Agent					
Concentration Operation			Į.		
Precipitation by Temperature Adjustment	0	0			0
Slurry Pulverization	1				
Slurry					
Water Content of Slurry [%]	45	45	45	45	45
Temperature of Slurry [°C]	40→6	40→7	40	70	40→7
	0	0			0
Increased Amount of	8.2	10.2	-	-	25.2
Undissolved Salt [%]					
Particle Properties					
Average Particle Size [µm]	248	245	260	244	238
Bulk Density [g/L]	608	615	598	620	614
Particle Strength [MPa]	30	30	28	30	32
Supporting Capacity [mL/g]	0.47	0.49	0.42	0.38	0.55
Mode Diameter of Microporous	1.2	1.1	1.9	1.6	0.95
Capacity Distribution [µm]		ĺ			
0.01 - 3 μm [mL/g]	0.33	0.32	0.37	0.32	0.31

Table 6

	Ex. 8	Ex. 9	Comp. Ex. 5	Comp. Ex. 6	Ex. 10
Composition of Detergent Parts by Weight					
Particles for Supporting Surfactant 12	100				
Particles for Supporting Surfactant 13		100			Ì
Particles for Supporting Surfactant 14			100		}
Particles for Supporting Surfactant 15				100	
Particles for Supporting Surfactant 16					100

Table 6 (continued)

	Ex. 8	Ex. 9	Comp. Ex. 5	Comp. Ex. 6	Ex. 10
Composition of Detergent Parts by Weight	1				
Surfactants	50	50	50	50	50
(Sodium Alkylbenzenesulfonate)	(21)	(21)	(21)	(21)	(21)
(Polyoxyethylene Alkyl Ether)	(21)	(21)	(21)	(21)	(21)
(Polyethylene Glycol)	(4)	(4)	(4)	(4)	(4)
(Water)	(4)	(4)	(4)	(4)	(4)
Amorphous	6	5	8	10	3
Aluminosilicate		:			-
Properties				_	
Average Particle Size [µm]	265	267	278	266	250
Bulk Density [g/L]	740	752	732	745	748
Flowability [s]	6.1	6.2	6.3	6.2	6.1
Bleed-out Property	1	1	1	1	1

Example 11

10

15

25

[0253] A first preparation liquid prepared in the same manner as in Comparative Example 1 was subjected to wet pulverization by COLLOID MILL, Model: MZ-80 (manufactured by SHINKO PANTEC CO., LTD.) at a flow rate of 800 kg/h.

(1024) The number of particles and the particle size distribution before and after the pulverization in the preparation illiquid were determined by TSUB-TEC M100. Incidentally, during the determination, in the same manner as in Example 4, there were provided alliquid corresponding to a first preparation includ prepared a separate mixing vessel without blending zeolite, and a liquid corresponding to a second preparation liquid prepared by pulverizing the liquid corresponding to a first preparation liquid was 778 counts/s, and the average particle size (on a number basis) was 172 µm. The number of particles in the liquid corresponding to a first preparation liquid was 778 counts/s, and the average particle size (on a number basis) was 172 µm. The number of particles of the liquid corresponding to a second preparation liquid after the pulverization was 2648 counts/s, and the average particle size was 24.6 µm. From these determination results, the number of particles of the water-soluble salt was increased by 2478 counts/s by the pulverization. The pulverizat second preparation liquid was spray-dried in the same manner as in Example 1. The high-temperature gas to be fed to the spray-driped flower was fed at a temperature of 200°C from the totom of the tower, and exhausted at 90°C from the top of the tower. The water content of the resulting Particles for Supporting Surfactant 17 was 4%.

[0255] Detergent Particles 17 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 17. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 5 parts by weight.

Example 12

[0256] A second preparation liquid prepared in the same manner as in Example 1 was subjected to wet pulverization by CAVITRON Model: CD1101 (manufactured by PACIFIC MACHINERY & ENGINEERING CO., LTD.) under the conditions of a rotational speed of 11200 pm at a flow rate of 800 kg/h.

[0257] The number of particles and the particle size distribution before and after the pulverization in the preparation liquid were determined by TSUB-TEC M100. Incidentally, the determination was carried out in the same manner as in Example 11. The number of particles in the liquid corresponding to a first preparation liquid was 778 counts/s, and the average particle size was 172 µm. The number of particles in the preparation liquid before the pulverization was 2634 counts/s, and the average particle size (on a number basis) was 21.2 µm. The number of particles in the liquid corresponding to a second preparation liquid after the pulverization was 4875 counts/s, and the average particle size was 18.4 µm. From these determination results, the number of particles of the water-soluble salt was increased by 2041 counts/s by the pulverization.

[0258] The pulverized second preparation liquid was spray-dried in the same manner as in Example 1. In addition, the particle constituting the resulting supporting particles was analyzed for a cave-in hole. As a result, the particles were composed of 85% of cave-in particles, in which a hole having a projected area diameter of 2 to 70% of a projected area diameter of a particle and a depth of 10% or more of the projected area diameter of the particle was present at

one or more points. In addition, the average value of

projected area diameter of hole projected area diameter of particle × 100

of a cave-in hole for the above 90% of cave-in particles was 15%.

[0259] Detergent Particles 18 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 15. The amount of an amorphous alluminositicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 5 parts by weight.

Example 13

10

[0260] A second preparation liquid having a water content of 45% by weight prepared in the same manner as in Example 5 was subjected to wet pulverization by COLLOID MILL. Model: MZ-80 at a flow rate of 800 kg/h.

[0261] The number of particles and the particle size distribution before and after the pulverization in the preparation liquid were determined by TUB-TEC M100. Incidentility, the determination was carried out before and after pulverizing a liquid corresponding to a second preparation liquid, which was prepared without blending zeolle in Example 5. The number of particles in the preparation liquid before the pulverization was 6351 counts/s, and the average particle size (on a number basis) was 20.0 µm. The number of particles in the liquid corresponding to a second preparation liquid after the pulverization was 8916 counts/s, and the average particle size was 17.0 µm. From these determination results, the number of particles of the water-soluble sait was increased by 2556 counts/s by the pulverization.

[0262] The pulverized second preparation liquid was spray-dried in the same manner as in Example 1. The hightemperature gas to be fed to the spray-drying tower was fed at a temperature of 220°C from the bottom of the tower, and exhausted at 110°C from the top of the tower. The water content of the resulting Particles for Supporting Surfactant 19 was 4%.

[0263] Detergent Particles 19 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 19. The amount of an amorphous aluminositicate supplied, as the minimum amount in which the bleed-out property of the deteroent particles is to be evaluated as 1, was 0.5 carts by weight.

Example 14

35

[0264] A second preparation liquid at 70°C prepared in the same manner as in Example 9 was subjected to wet pulverization by CAVITRON Model: CD1010 under the conditions of a rotational speed of 11200 rpm at a flow rate of 800 kg/h.

[0265] The number of particles and the particle size distribution before and after the pulverization in the preparation liquid were determined by TUB-TEC M100. Incidentally, the determination was carried out before and after pulverizing a liquid corresponding to a second preparation liquid, which was prepared without formulating zealits in Example 9. The number of particles in the preparation liquid before the pulverization was 8255 counts/s, and the average particle size (on a number bass) was 28.0 µm. The number of particles in the liquid corresponding to a second preparation liquid after the pulverization was 11831 counts/s, and the average particle size was 20.3 µm. From these determination results, the number of particles of the water-soluble salt was increased by 3576 counts/s by the pulverization. The pulverized second preparation liquid was spray-dried in the same manner as in Example 1. The high-temperature gas to be fed to the spray-drying tower was fed at a temperature of 220°C from the bottom of the tower, and exhausted at 110°C from the top of the tower. The water content of the resultin Particles for Supporting Surficant 20 was 49°C.

[0266] Detergent Particles 20 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 20. The amount of an amorphous aluminosilicate supplied, set the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 3.5 parts by weight.

[0267] The composition, the properties and the like of each group of the resulting Particles for Supporting Suffactant 17 to 20 are shown in Table 3, and the properties of each group of Detergent Particles 1 to 20 are shown in Table 8. [0268] As shown in the results of Tables 7 and 8, by subjecting the particles of the water-soluble salt in a sturry to wet pulverization to increase the number of the particles, the supporting ability of the particles for supporting a surfactant could be improved, and the amount of the amorphous aluminosilizate could be reduced. In addition, the more the amount of undissolved substance in a sturry, the greater the effect of the improvement in the supporting ability of the particles for supporting a surfactant by wet pulverization.

Table 7

	Ex. 11	Ex. 12	Ex. 13	Ex. 14
Composition % By Weight				
Zeolite	44.0	44.0	44.0	44.0
Sodium Carbonate	21.0	21.0	23.0	23.0
Sodium Sulfate	21.0	21.0	23.0	23.0
Sodium Sulfite	8.0	0.8	0.8	0.8
Sodium Polyacrylate	5.0	5.0	5.0	5.0
Fluorescent Dye	0.2	0.2	0.2	0.2
Sodium Chloride	4.0	4.0	0.0	0.0
Sodium Bromide	0.0	0.0	0.0	0.0
Water	4.0	4.0	4.0	4.0
TOTAL	100.0	100.0	100.0	100.0
Operation			-	
Post-Addition of Microcrystal-		0		
Precipitating Agent				
Concentration Operation			0	
Precipitation by Temperature	- 1			0
Adjustment	- 1			
Slurry Pulverization	0	0	0	0
Slurry				
Water Content of Slurry [%]	42	42	50>45	45
Temperature of Slurry [°C]	40	40	60	40→70
Increased Amount of	-	16.3	19	10.2
Undissolved Salt [%]	l			
Particle Properties		i		
Average Particle Size [µm]	240	252	258	244
Bulk Density [g/L]	604	605	602	610
Particle Strength [MPa]	31	31	30	30
Supporting Capacity [mL/g]	0.42	0.51	0.65	0.54
Mode Diameter of Microporous	1.05	0.76	0.56	0.92
Capacity Distribution [µm]	- 1	- 1		
0.01 - 3 μm [mL/g]	0.3	0.32	0.36	0.32

Table 8

£x. 14
100
50
(21)
(21)
(4)
(4)

Table 8 (continued)

	Ex. 11	Ex. 12	Ex. 13	Ex. 14
Composition of Detergent Parts by Weight				
Amorphous Aluminosilicate	8	5	0.5	3.5
Properties				
Average Particle Size [µm]	251	267	273	256
Bulk Density [g/L]	743	741	750	755
Flowability [s]	6.2	6.2	6.1	6.3
Bleed-out Property	1	1	1	1

15 Example 15

10

[0269] A first preparation liquid having a water content of 51% by weight was prepared in the same manner as in Example 4, and subjected to we pluvefization by COLLOID MILL, Model: M2.8 da a flow rate of 800 kg/h. Thereafter, the ground first preparation liquid was subjected up to a concentration operation to a water content of 48% by weight, to give a second preparation liquid was subjected up to a concentration operation to a water content of 48% by weight, to give a second preparation liquid was spray-dried, to give Particles for Supporting Surfactant 21. Detergent Particles for Supporting Surfactant 21. The amount of an amorphous aluminosilicate fed, as the minimum amount in which the bleed-out property of the deterent particles is to be evaluated as 1, was 7 parts by weight.

25 Comparative Example 7

[0270] A first preparation liquid having a water content of 48% by weight was prepared in the same manner as in Example 15, and spray-dried without carrying out wet pulverization and concentration, to give Particles for Supporting Surfactant 22. Detergent Particles 22 were prepared in the same manner as in Example 15 using Particles for Supporting Surfactant 22. However, in the case where the amorphous aluminosilicate was used in an amount of 7 parts by weight, the same amount as that of Example 15, Particles for Supporting Surfactant 22 did not sufficiently support the surfactant composition during the agitation in a Lödige Mixer, and became aggregated, so that the values of the properties were deteriorated to an extent to be undeterminable.

35 Example 16

[0271] A first preparation liquid having a water content of 48% by weight was prepared in the same manner as in Example 8, and subjected to wet pulvetization by COLLOID MILL, Model: M2-60 at a flow rate of 800 kg/h. Thereafter, the preparation liquid was heated to 70°C, to give a second preparation liquid. This second preparation liquid was spray-dried, to give Particles for Supporting Surfactant 23. Detergent Particles 23 am expression as meanment as in Example 1 using Particles for Supporting Surfactant 23. The amount of an amorphous aluminositicate supplied, as the minimum amount in which the bleed-out property of the detergent particles is to be evaluated as 1, was 7 parts by weight.

45 Comparative Example 8

[0272] A first preparation liquid having a water content of 48% by weight was prepared in the same manner as in Example 16, and spray-dried without carrying out wet pulverization and concentration, to give Particles for Supporting Surfactant 24. Detergent Particles for Supporting Surfactant 24. However, in the case where the amorphous aluminosilicate was formulated in an amount of 7 parts by weight, the same amount as that of Example 16, Detample 16, Deta

[0273] The composition, the properties and the like of each group of the resulting Particles for Supporting Surfactant 21 to 24 are shown in Table 9, and the properties of each group of Detergent Particles 21 to 24 are shown in Tables 0. [0274] From the results shown in Tables 9 and 10, the supporting ability of the particles for supporting a surfactant was improved, even when the concentration and the heating operation were carried out after subjecting the first preparation liquid to wet outwersization.

Table 9

35 .

	Ex. 15	Comp. Ex. 7	Ex. 16	Comp. Ex. 8
Composition % By Weight	1			
Zeolite	40.0	40.0	40.0	40.0
Sodium Carbonate	13.0	13.0	36.0	36.0
Sodium Sulfate	36.0	36.0	13.0	13.0
Sodium Sulfite	0.8	0.8	0.8	0.8
Sodium Polyacrylate	6.0	6.0	6.0	6.0
Fluorescent Dye	0.2	0.2	0.2	0.2
Sodium Chloride	0.0	0.0	0.0	0.0
Sodium Bromide	0.0	0.0	0.0	0.0
Water	4.0	4.0	4.0	4.0
TOTAL	100.0	100.0	100.0	100.0
Operation				
Post-Addition of Microcrystal-	0			
Precipitating Agent	1			
Concentration Operation				
Precipitation by Temperature	1		0	
Adjustment				
Siurry Pulverization	0		0	
Slurry				-
Water Content of Slurry [%]	51→48	48	48	48
Temperature of Slurry [°C]	50	50	40→70	40
Increased Amount of	11.6	-	9	
Undissolved Salt [%]				
Particle Properties				
Average Particle Size [µm]	225	205	210	198
Bulk Density [g/L]	545	551	505	460
Particle Strength [MPa]	22	16	17	12
Supporting Capacity [mL/g]	0.45	0.38	0.46	0.4
Mode Diameter of Microporous	1.12	1.89	1.2	1.5
Capacity Distribution [µm]				
0.01 - 3 μm [mL/g]	0.33	0.28	0.36	0.38

Table 10

	lable 10			
	Ex. 15	Comp. Ex. 7	Ex. 16	Comp. Ex. 8
Composition of Detergent Parts by Weight	-			
Particles for Supporting Surfactant 21 Particles for Supporting Surfactant 22 Particles for Supporting Surfactant 23 Particles for Supporting Surfactant 24	100	100	100	100
Surfactants	50	50	50	50
(Sodium Alkylbenzenesulfonate)	(21)	(21)	(21)	(21)
(Polyoxyethylene Alkyl Ether)	(21)	(21)	(21)	(21)
(Polyethylene Glycol)	(4)	(4)	(4)	(4)
(Water)	(4)	(4)	(4)	(4)

Table 10 (continued)

	Ex. 15	Comp. Ex. 7	Ex. 16	Comp. Ex. 8
Composition of Detergent Parts by Weight				
Amorphous Aluminosilicate	7	7	7	7
Properties				
Average Particle Size [µm]	243	Undeterminable	231	240
Bulk Density [g/L]	721	Undeterminable	702	623
Flowability [s]	6.2	Undeterminable	6.2	8.6
Bleed-out Property	1 :	5	1	4

15 Example 17

5

10

26

[0275] Particles for Supporting Surfactant 25 were obtained in the same manner as in Example 12. Incidentally, as a 40% by weight aqueous solution of sodium polyacrylate, there was used one prepared according to the following method.

[0276] An amount 8.0.3 kg of water was supplied, and heated to 100°C. While keeping the temperature at 100°C, 190 kg (2.1 kmol) of 80% by weight acrylic acid and 3.0 kg (4.6 kmol) of a 98% aqueous solution of 2-mercaptoethanol are added dropwise at a constant rate over 4 hours, and 5.0 kg (6.3 md) of a 30% by weight aqueous sodium persulfate is added dropwise at a constant rate over 6 hours, to carry out polymerization. After the termination of the dropping polymerization, 2.1 kg (217.6 kmol) of a 35% by weight aqueous solution of hydrogen persoxide is added dropwise over 1 hour for deodorization. Further, the resulting mixture is matured for 4 hours, and cooled. When the internal temperature is 60°C, 3.3 kg (11.5 mol) of a 35% by weight aqueous sodium hydrogensultie is added as a reducing agent, and the resulting mixture is reacted for 1 hour. Thereafter, the mixture was cooled, and 167 kg (2 kmol) of a 48% by weight aqueous sodium hydroxide was added thereto, while keeping the temperature 40°C or lower. Water was added the reto, while keeping the temperature 40°C or lower. Water was added the reto. While keeping the temperature 40°C or lower. Water was added the reto. While keeping the temperature 40°C or lower. Water was added the reto. 1000 to 10

[0277] Method for Molecular Weight Determination

- 1. Standard substance for calculation: polyacrylic acid (AMERICAN STANDARDS CORP)
- 2. Eluent: 0.2 mol/L phosphate buffer/CH3CN: 9/1 (volume ratio)
- 3. Column: PWXL + G4000PWXL + G2500PWXL (manufactured by Tosoh Corporation)
- 4. Detector: RI
 - 5. Sample concentration; 5 mg/mL
- 6. Injected amount: 0.1 mL
- 7. Temperature for determination: 40°C
 - 8. Flow rate: 1.0 mL/min

[0278] In addition, the particle constituting the resulting supporting particles was analyzed for a cave-in hole. As a result, the particles were composed of 90% of cave-in particles, in which a hole having a projected area diameter of 2 to 70% of a projected area diameter of a particle and a depth of 10% or more of the projected area diameter of the particle was present at one or more points, in addition, the average value of

$\frac{\text{projected area diameter of hole}}{\text{projected area diameter of particle}} \times 100$

of a cave-in hole for the above 90% of cave-in particles was 19%.

[0279] Detergent Particles 21 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 25. Detergent Particles 25 had a sufficiently excellent flowability, and the level of the bleed-out property was evaluated as 1 without addition of an amorphous aluminositicate.

55 Example 18

[0280] A first preparation liquid having a water content of 55% by weight was prepared in the same manner as in Example 6, and thereafter the first preparation liquid was subjected to a concentration operation up to a water content

of 51% by weight. Further, a microcrystal-precipitating agent was added to adjust the preparation liquid to a water content of 50% by weight, and thereafter the resulting preparation liquid was spray-dried, to give Particles for Supporting Surfactant 26. Detergent Particles 26 were prepared in the same manner as in Example 1 using Particles for Supporting Surfactant 26. At this time, 55 parts by weight to a surfactant composition were supplied.

[0281] Detergent Particles 26 had a sufficiently excellent flowability, and the level of the bleed-out property was evaluated as 1 without addition of an amorphous aluminosilicate.

Comparative Example 9

25

30

35

40

45

50

[0282] Particles for Supporting Surfactant 27 were obtained in the same manner as in Comparative Example 1. Detergent Particles 27 were prepared in the same manner as in Example 17 using the resulting Particles for Supporting Surfactant 27. An amorphous aluminositicate was not added in the same manner as in Example 17. However, since the supporting abulls of Particles for Supporting Surfactant 27 is lower than that of Particles for Supporting Surfactant 27. Barticles for Supporting Surfactant 27 is provided for Supporting Surfactant composition and became aggre-

gated in a Lödige Mixer, so that the values of the properties were deteriorated to an extent to be undeterminable. [0283] The composition and the properties of each group of the resulting Particles for Supporting Surfact

can be further improved depending on the composition of particles for supporting a surfactant or the water content of the preparation liquid. Since each group of Particles for Supporting Surfactant 25 and 26 obtained according to the method of the present invention has a mode diameter of micropports capacity distribution of 1.5 mm or less and a high supporting ability, detergent particles having an excellent bleed-out property can be obtained by using these groups of particles without addition of an amorphous aluminosilicate, and a higher amount of the surfactant composition could be further formulated.

Table 11

181	ne II		
	Ex. 17	Ex. 18	Comp. Ex. 9
Composition % By Weight			İ
Zeolite	36.0	36.0	36.0
Sodium Carbonate	25.0	25.0	25.0
Sodium Sulfate	23.8	23.8	23.8
Sodium Sulfite	1.0	1.0	1.0
Sodium Polyacrylate	6.0	6.0	5.0
Fluorescent Dye	0.2	0.2	0.2
Sodium Chloride	4.0	4.0	4.0
Sodium Bromide	0.0	0.0	0.0
Water	4.0	4.0	4.0
TOTAL	100.0	100.0	100.0
Operation			
Post-Addition of Microcrystal-	0	0	
Precipitating Agent			
Concentration Operation		0	
Precipitation by Temperature			
Adjustment			
Slurry Pulverization	0		
Siurry			
Water Content of Slurry [%]	50	55→50	50
Temperature of Slurry [°C]	50	60	50
Increased Amount of	11.7	24.1	-
Undissolved Salt [%]			
Particle Properties			
Average Particle Size [µm]	259	263	250

Table 11 (continued)

	Ex. 17	Ex. 18	Comp. Ex. 9
Particle Properties			
Bulk Density [g/L]	542	548	579
Particle Strength [MPa]	30	36	15
Supporting Capacity [mL/g]	0.68	0.72	0.52
Mode Diameter of Microporous	0.81	0.48	1.63
Capacity Distribution [µm]			
0.01 - 3 μm [mL/g]	0.43	0.45	0.5

Table 12

lable 1	2		
	Ex. 17	Ex. 18	Comp. Ex. 9
Composition of Detergent Parts by Weight		Ċ	
Particles for Supporting Surfactant 25	100		
Particles for Supporting Surfactant 26		100	
Particles for Supporting Surfactant 27			100
Surfactants	50	55	50
(Sodium Alkylbenzenesulfonate)	(21)	(23)	(21)
(Polyoxyethylene Alkyl Ether)	(21)	(23)	(21)
(Polyethylene Glycol)	(4)	(4.5)	(4)
(Water)	(4)	(4.5)	(4)
Amorphous Aluminosilicate	0	0	5
Properties			
Average Particle Size [μm]	272	274	Undeterminable
Bulk Density [g/L]	738	743	Undeterminable
Flowability [s]	6.2	6.1	Undeterminable
Bleed-out Property	1	1	5

Example 19

5

10

15

30

35

[0.285] A mixing vessel equipped with a jacket, comprising an agitator, was charged with 550 parts by weight of water. After the water temperature reached 35°C, 72 parts by weight of sodium carbonate, 194 parts by weight of sodium sulfate, and 83 parts by weight of a 40% by weight aqueous solution of sodium polyacrylate were sequentially added thereto. The resulting mixture was agitated for 30 minutes, to give a homogenous aqueous solution in which watersoluble components were completely dissolved (water content; 70% by weight).

[0286] The aqueous solution was spray-dried in the same manner as in Example 1. The high-temperature gas to be fed to the spray-drying tower was fed at a temperature of 230°C from the bottom of the tower, and exhausted at 95°C from the top of the tower. The water content of the resulting particle was 5% by weight.

[0287] The particle was subjected to dry pulverization using ATOMIZER, Model: EIIW-7.5 (manufacturer by Fuji Paudal Co., Lid.) under the conditions of the climenter of a screen mesh of 0.5 mm, a feed amount for pulverization of 60 kg/h; and a rotational speed of 5000 rpm, to give a fine powder having an average particle size of 5 μm (hereinafter referred to as a fine powder).

[0288] In addition, another mixing vessel equipped with a jacket, comprising an agitator, was charged with 452 parts by weight of water. After the water temperature reached 35°C, 65 parts by weight of sodium sulfite, parts by weight of sodium sulfite, and 1 part by weight of a fluorescent dye were added thereto, and the resulting mixture was agitated for 0 minutes. One-hundred and twenty-three parts by weight of sodium carbonate were added to mixture, and 64 parts by weight of a e05°K by weight and could une polyacybate were added thereto. The resulting mixture was agitated for 10 minutes. Further, 1980 parts by weight of 2 mixture were added and the resulting mixture was agitated for 30 minutes. Further, 1980 parts by weight of 2 mixture added thereto, and the resulting mixture was agitated for 30 minutes, to give a second preparation liquid (water content: 50% by weight). The final temperature of this second preparation liquid water content: 50% by weight).

[0289] After the preparation of the first preparation liquid and 10 minutes after the addition of the fine powder, a sample was taken from each of the preparation liquids, and the number of particles and the particle size distribution were determined by TSUB-TEC M100. In the first preparation liquid, an inorganic stall was entirely dissolved, so that the number of particles was hardly detected. The number of particles in the second preparation liquid after the addition of the fine powder was 4009 countists, and the average particle size was 10.5 µm.

[0290] The second preparation liquid was spray-dried in the same manner as in Example 1. The high-temperature gas to be fed to the spray-drying tower was fed at a temperature of 220°C from the bottom of the tower, and exhausted at 110°C from the top of the tower. The water content of the resulting Particles for Supporting Surfactant 28 was 4% by weight.

[0231] Detergent Particles 28 were prepared in the same manner as in Example 1 using the resulting Particles for Supporting Surfactant 28. The amount of an amorphous aluminosilicate supplied, as the minimum amount in which the biled-out property of the detergent particles is to be evaluated as 1, was 3 parts by weight.

Comparative Example 10

15

25

35

40

45

50

55

[0292] Particles for Supporting Surfactant 29 were obtained in the same manner as in Example 19 except that a fine particle was not added. Detergent Particles 29 were prepared in the same manner as in Example 19 using the resulting Particles for Supporting Surfactant 29. However, when the amorphous aluminosities was added in an amount of 3 parts by weight, the same amount as that of Example 19, the particles for supporting a surfactant did not sufficiently support the surfactant composition and became aggregated in a Lödige Mixer, so that the values of the properties were deteriorated to an extent to be undeterminable.

ICES31 The composition and the properties of each group of the resulting Particles for Supporting Surfactant 28 and 29 are shown in Table 13, and the properties of each group of Detergent Particles 28 and 29 are shown in Table 14. ICES41 Particles for Supporting Surfactant 29 of Comparative Example 10, in which a fine particle of a water-scholause sails not added, have a poor particle strength and a large mode diameter of microporous capacity distribution. Therefore, bleeding-out of a surfactant composition, which was once absorbed in the particles for supporting a surfactant, due to disintegration of the particle, and the like, was found in the step of supporting the surfactant composition, such that the properties of the detergent particles were drastically deteriorated. On the other hand, since Particles for Supporting Surfactant 28 have a relatively high particle strength while having the same composition, and has a mode diameter of microporous capacity distribution of 1.5 µm or less and a high supporting ability, the amorphous aluminositicate used was considerably reduced when using Particles for Supporting Surfactant 28.

Table 13

Table to		
	Ex. 19	Comp. Ex. 10
Composition % By Weight		
Zeolite	38.0	38.0
Sodium Carbonate	26.0	26.0
Sodium Sulfate	24.8	24.8
Sodium Sulfite	1.0	1.0
Sodium Polyacrylate	6.0	6.0
Fluorescent Dye	0.2	0.2
Sodium Chloride	0.0	0.0
Sodium Bromide	0.0	0.0
Water	4.0	4.0
TOTAL	100.0	100.0
Operation		
Post-Addition of Microcrystal-		
Precipitating Agent		
Concentration Operation		
Increased Amount of Undissolved Salt [%]		
Slurry Pulverization		
Addition of Fine Particle	0	

Table 13 (continued)

	Ex. 19	Comp. Ex. 10
Slurry		
Water Content of Slurry [%]	50	50
Temperature of Slurry [°C]	50	50
Increased Amount of	23.3	-
Undissolved Salt [%]		
Particle Properties		
Average Particle Size [µm]	255	269
Bulk Density [g/L]	510	461
Particle Strength [MPa]	25	12
Supporting Capacity [mL/g]	0.57	0.44
Mode Diameter of Microporous	0.88	1.85
Capacity Distribution [µm]		
0.01 - 3 μm [mL/g]	0.48	0.4

Table 14

	Ex. 19	Comp. Ex. 10
Composition of Detergent Parts by Weight	1	
Particles for Supporting Surfactant 28	100	
Particles for Supporting Surfactant 29		100
Surfactants	50	50
(Sodium Alkylbenzenesulfonate)	(21)	(21)
(Polyoxyethylene Alkyl Ether)	(21)	(21)
(Polyethylene Glycol)	(4)	(4)
(Water)	(4)	(4)
Amorphous	3	8
Aluminosilicate		•
Properties		
Average Particle Size [µm]	270	Undeterminable
Bulk Density [g/L]	723	Undeterminable
Flowability [s]	6.3	Undeterminable
Bleed-out Property	1	5

Example 20

10

15

25

30

35

[0285] A mixing vessel was charged with 430 parts by weight of water. After the water temperature reached 35°C, 108 parts by weight of sodium suffate, 5 parts by weight of sodium suffate, and 2 parts by weight of a fluorescent dye were added thereto, and the resulting mixture was agitated for 10 minutes. One-hundred and fifteen parts by weight of sodium carbonate were added to the mixture, and 150 parts by weight of a 40% by weight aqueous solition of sodium polyacrylate were added thereto. The resulting mixture was agitated for 10 minutes, to give a first preparation liquid. Forty parts by weight of sodium chloride, which was a microcrystal-precipitating agent, were added thereto, and the resulting mixture was agitated for 10 minutes. Subsequently, the mixture was subjected to wet pulverization by COLLOID MILL, Model: M2-80 at a flow rate of 800 kg/h. Thereafter, 150 parts by weight of zerollite were added, and the resulting mixture was agitated for 30 minutes, to give a homogenous second preparation liquid (water content of surry; 52% by weight). The final temperature of this preparation liquid was 50°C. The amount of the water-soluble inorganic salt precipitated by the addition of sodium chloride was 17.8% by weight of that dissolved in the first preparation liquid was 50°C.

[0296] After the preparation of the first preparation liquid, 10 minutes after the addition of sodium chloride, and after

the puterization of the preparation liquid, a sample was taken from each of the preparation liquids, and the number of particles and the partiel size distribution were determined by TSUB-TEC M100. The number of particles in the first preparation liquid was 557 counts/s, and the average particle size (on a number basis) was 125 µm. The number of particles in the preparation liquid after the addition of sodium chloride was 3796 counts/s, and the average particle size was 20.3 µm. From these cetermination results, the number of microcystals was increased by 3241 counts/s by the addition of sodium chloride, and the average particle size of the increased microcrystals was 17.0 µm. In addition, the number of particles in the second preparation liquid after the puterization was 5438 counts/s, and the average particle size was 18.2 µm. The number of particles of the water-soluble salt was additionally increased by 1540 counts/s by the outlerization.

[0237] Spray-drying was carried out in the same manner as in Example 12, to give Particles for Supporting Surfactant 30. Delargent Particles 30 were prepared using Particles for Supporting Surfactant 30 by the method shown below.
[0238] A surfactant composition (polyosythylene alkyl etherholyethylene glycol/sodium alkylbenzenesul/onatelwater = 25/5/25/5 (weight ratio)) was adjusted to 80°C. Next, 100 parts by weight of the resulting particles for supporting a surfactant were supplied into a Lödige Mixer (manufactured by Matsuzaka Giken Co., Ltd.; capacity 130 L; equipped with a jacket), and the agitation of a main shaft (agitation impellers; rotational speed: 60 rpm; peripheral speed: 1.6 m/ s) was started incidentally, hot water at 80°C was allowed to flow through the jacket at 10 L/minute. Suity parts by weight of the above surfactant composition were supplied into the above mixer lar minutes, and thereafter the resulting mixture was agisted for 5 minutes. Further, 20 parts by weight of a crystalline silicate, and zeolite were supplied thereinto. The agitations of the main shaft (orgational speed: 120 rpm; peripheral speed:

3.1 m/s) and a chopper (rotational speed: 3600 rpm; peripheral speed: 28 m/s) were carried out for 1 minute, and Detergent Particles 30 were discharged. The minimum amount of zeolite in which the bleed-out property of the detergent particles is to be evaluated as 1 was 3 parts by weight.

Example 21

25

40

55

[0299] Particles for Supporting Surfactant 31 were obtained in the same manner as in Example 20 except that a 40%, by weight acqueous solution of sodium polyacrylate was supplied together with water when a first preparation licuid was prepared. Detergent Particles 31 were prepared in the same manner as in Example 20 using the resulting Particles for Supporting Surfactant 31. Incidentally, Detergent Particles 31 had a sufficiently excellent flowability, and the level of the bleed-out property was evaluated as 1 without addition of scotile.

Comparative Example 11

[0300] Particles for Supporting Surfactant 32 were obtained in the same manner as in Example 1 except that a microcrystal-precipitating agent was not added. Detergent Particles 32 were prepared in the same manner as in Example 20 using the resulting Particles for Supporting Surfactant 32. The minimum amount of zeolite in which the bleed-out property of the detergent particles is to be evaluated as 1 was 16 parts by weight.

Comparative Example 12

[0301] Particles for Supporting Surfactant 33 were obtained in the same manner as in Comparative Example 11 except that a 40% by weight aqueous solution of sodium polyacrylate was supplied together with water when a first preparation liquid was prepared. Detergent Particles 33 were prepared in the same manner as in Example 20 using the resulting Particles for Supporting Surfactant 33. The minimum amount of zeolite in which the bleed-out property of the detergent particles is to be evaluated as 1 was 13 parts by weight.

[9302] The composition and the properties of each group of the resulting Particles for Supporting Surfactant 30 to 33 are shown in Table 15, and the properties of each group of Detergent Particles 30 to 33 are shown in Table 16, and the properties of each group of Detergent Particles 30 to 33 are shown in Table 16, illy of the particles for supporting a surfactant was improved, and the amount of zeolite for surface-modifying could be dramatically reduced, even in the case where the amount of the polymer formulated was increased. In addition, at the time of preparing the first preparation liquid, when the water-soluble polymer was added prior to adding of sodium carbonate, the supporting ability of the particles for supporting a surfactant was improved. However, its effect was small, as compared to the effect of the improvement in the supporting ability by the technique according to the present invention.

Table 15

iaui	6 13		
Ex. 20	Ex. 21	Comp. Ex. 11	Comp. Ex. 12
30.0	30.0	30.0	30.0
23.0	23.0	27.0	27.0
21.6	21.6	25.6	25.6
1.0	1.0	1.0	1.0
12.0	12.0	12.0	12.0
0.4	0.4	0.4	0.4
8.0	8.0	0.0	0.0
0.0	0.0	0.0	0.0
4.0	4.0	4.0	4.0
100.0	100.0	100.0	100.0
			-
0	0		
1			
-			
0	0		-
I			
52	52	52	52
50	50	50	50
17.8	18.6	-	-
255	248	244	243
536	525	503	512
35	35	21	23
0.62	0.68	0.52	0.54
0.72	0.68	2.20	1.80
1			
0.49	0.49	0.47	0.47
	30.0 21.6 1.0 23.0 24.6 1.0 0.4 4.0 0.0 4.0 0.0 100.0 0 0 17.8 255 536 0.62 0.72	30.0 30.0 30.0 23.0 23.0 23.0 23.0 23.0	Ex. 20 Ex. 21 Comp. Ex. 11 30.0 30.0 30.0 30.0 23.0 23.0 27.0 21.6 21.6 21.6 25.6 1.0 1.0 1.0 12.0 4.0 4.0 4.0 4.0 8.0 8.0 0.0 0.0 0.0 0.0 4.0 4.0 4.0 4.0 100.0 100.0 100.0 O C 52 52 52 52 50 50 50 17.8 18.6 - 255 248 244 536 525 503 35 35 21 0.62 0.68 0.52 0.72 0.68 0.52

Table 16

Table 10			
Ex. 20	Ex. 21	Comp. Ex. 11	Comp. Ex. 12
1			
100			
	100		
		100	
			100
60	60	60	60
(25)	(25)	(25)	(25)
(25)	(25)	(25)	(25)
(5)	(5)	(5)	(5)
(5)	(5)	(5)	(5)
	60 (25) (25) (5)	Ex. 20 Ex. 21 100 100 60 60 (25) (25) (25) (25) (5) (5)	Ex. 20 Ex. 21 Comp. Ex. 11 100 100 100 60 60 60 60 (25) (25) (25) (25) (25) (25) (25) (25) (5) (5) (5) (5)

Table 16 (continued)

	Ex. 20	Ex. 21	Comp. Ex. 11	Comp. Ex. 12
Composition of Detergent Parts by Weight				
Zeolite	3	0	15	12
Crystalline Silicate	20	20	20	20
Properties				
Average Particle Size [µm]	263	255	261	257
Bulk Density [g/L]	716	725	680	694
Flowability [s]	6	6.1	6.5	6.4
Bleed-out Property	1	1	1	1

Example 22

10

20

[0304] Particles for Supporting Surfactant 34 were obtained in the same manner as in Example 1. Incidentally, as a 40% by weight a queues solution of sodium polycyrdate, there was used one prepared according to the method described in Examples of Japanese Examined Patent Publication No. Hel 2-24283. The reaction was carried out by supplying an equeues solution of sodium acrylate having a neutralization degree of 95% and a concentration of 37.7% by weight at a rate of 3.11 kg/h, and supplying an equeues solution of sodium hydrogensulfite having a concentration of 35% by weight at a rate of 0.13 kg/h, at an average temperature of the jacket of 20°C with an air feeding rate of 3 m3/h. The weight-average molecular weight was 10000. In addition, the particle constituting the resulting particles for supporting a surfactant was analyzed for a cave-in hole. As a result, the particles were composed of 91% of cave-in particles, in which a hole having a projected area diameter of 2 to 75% of a projected area diameter of a particle and a depth of 10% or more of the projected area diameter of a particle and a depth of 10% or more of the projected area diameter of the particle was present at one or more points. In addition, the average value of

projected area diameter of hole projected area diameter of particle × 100

of a cave-in hole for the above 91% of cave-in particles was 17%. Also, the average value for the depth of the cavein hole was 55% of the projected area dismeter of the particle. The composition and the values of the properties of the resulting particles for supporting a surfactant are shown in Table 17. Incidentally, the absorbency of the ingular surfactant composition determined by the above-described method was expressed as a great value of 0.45 mL/g, so that the liquid surfactant composition was excellent in the absorbency.

Comparative Example 13

- [0305] Particles for Supporting Surfactant 35 were obtained in the same manner as in Comparative Example 3, Incidentally, "NEOPELEX F-65" (manufactured by Kao Corporation) was used as a 50% by weight aqueous solution of sodium alkylenzenesupionate, in the first preparation liquid, which was used for spray-drying, the water-soluble salt was completely dissolved. In addition, the particle constituting the resulting supporting particles was analyzed for a cave-in hole. As a result, there were substantially no cave-in particles, in which a hole having a projected area diameter of 2 to 70% of a projected area diameter of a particle and a depth of 10% or more of the projected area diameter of the particle was present at one or more points. The composition and the values of the properties of the resulting particles for supporting a surfactant are itself in Table 17, incidentally, the absorbency of the liquid surfactant composition determined by the above-described method was as small as 0.10 mL/g, indicating that the liquid surfactant composition was poor in the absorbency.
- [0306] Each group of Detergent Particles 34 and 35 was obtained by adding a surfactant to each group of Particles for Supporting Surfactant 34 and 35 of Example ?2 and Comparative Example 13 at a ratio shown in Table 18, to support the surfactant thereby. To 10 parts by weight of polyoxyethylene allyle their under mixing at 80 °C, 1 24 by weight of polyethylene glycol, paimitic acid (LUNAC P-95, manufactured by Kao Corporation) corresponding to 0.7% by weight of sodium paimitale, a precursor of an alkylbenzenesulfonic acid (NEOPELEX GS, manufactured by Kao Corporation) corresponding to 12 parts by weight of a sodium alkylbenzenesulfonate, and an acqueous sodium hydroxide as a neutralizing agent were added, thereby preparing a hydrated surfactant composition having the composition shown in Table 18. Next, 50 parts by weight of the above base particles were supplied into a Lodigle Mixer (manufacture).

by Matsuzaka Giken Co., Ltd.; capacity: 20 L; equipped with a jacket), and the agitations of a main shaft (150 rpm) and a chopper (4000 rpm) were started. Incidentally, hot water at 80°C was allowed to flow through the jacket at 10 L/ minute. The above hydrated surfactant composition was supplied into the above mixer in 2 minutes, and thereafter the resulting mixture was agitated for 4 minutes. Subsequently, 10 parts by weight of a crystalline silicate and 10 parts by weight of zeolite were added to the mixture, and a 2-minute surface-coating operation was carried out, thereby giving each group of Detergent Particles 34 and 35. Further, 2 parts by weight of zeolite and 1% by weight of an enzyme granule were added, to give a granular detergent composition. The composition and the properties of the resulting detergent compositions are shown in Table 18. The detergent composition prepared using Particles for Supporting Surfactant 34 of Example 22 showed satisfactory values for the properties. On the other hand, in the case where Particles for Supporting Surfactant 35 of Comparative Example 13 were used, Particles for Supporting Surfactant 35 did not sufficiently support the surfactant composition within the time of the above operation and became aggregated. so that the values of the properties were deteriorated to an extent to be undeterminable.

15

25

30

35 -

45

50

Table 1	7	
	Ex. 22	Comp. Ex. 13
Composition % By Weight	1	
Zeolite	27.4	50.0
Sodium Carbonate	25.6	20.0
Sodium Sulfate	21.6	10.0
Sodium Sulfite	1.0	1.5
Sodium Polyacrylate	13.0	9.0
Fluorescent Dye	0.4	0.5
Sodium Chloride	8.0	0.0
Sodium Alkylbenzenesulfonate	0.0	4.0
Water	3.0	5.0
TOTAL	100.0	100.0
Operation		
Post-Addition of Microcrystal-		
Precipitating Agent	0	
Concentration Operation		
Increased Amount of		
Undissolved Salt [%]		
Slurry Pulverization	l .	
Slurry		
Water Content of Slurry [%]	53	50
Temperature of Slurry [°C]	50	58
Increased Amount of	21.5	-
Undissolved Salt [%]		
Particle Properties		
Average Particle Size [µm]	246	225
Bulk Density [g/L]	510	620
Particle Strength [MPa]	40	25
Cave-In Granule Ratio [%]	91	0
Average Diameter of	17	-
Cave-In Hole [%]		
Average Depth of	55	-
Cave-In Hole [%]		
Supporting Capacity [mL/g]	0.60	0.52
Absorbency [mL/g]	0.45	0.10
Mode Diameter of Microporous Capacity Distribution [µm]	0.73	1.60
Capacity Distribution [http]		

Table 17 (continued)

	Ex. 22	Comp. Ex. 13
Particle Properties		
0.01 - 3 μm [mL/g]	0.48	0.28

Toble 1

Ex. 22	Comp. Ex. 13
1	
50	
	50
27	27
(12)	(12)
(10)	(10)
(0.7)	(0.7)
(1.2)	(1.2)
(3.1)	(3.1)
12	12
10	10
1	1
275	Undeterminable
745	Undeterminable
6.2	Undeterminable
1	-
	27 (12) (10) (0.7) (1.2) (3.1) 12 10 1

· INDUSTRIAL APPLICABILITY

- 36 (9307) According to the present invention, there can be obtained particles for supporting a surfactant having excellent supporting ability (supporting capacity/supporting strength) of the liquid surfactant composition, runting excellent absorbersy (supporting rate) of the liquid surfactant composition. Further by supporting the surfactant having excellent absorbersy (supporting rate) of the surfactant composition to the particles for supporting a surfactant, detergent particles having excellent detergency performance, quality and the like can be efficiently obtained.
- (a) (2008) The present invention being thus described, it will be obvious that the same may be varied in many ways. Such variations are not to be regarded as a departure from the spirit and scope of the invention, and all such modifications as would be obvious to one skilled in the attare intended to be included within the scope of the following colors.

45 Claims

50

10

15

20

25

- 1. A process for proparing particles for supporting a surfactant comprising the steps of preparing a preparation liquid comprising a water-soluble polymer and a water-soluble salt, and spray-drying the preparation liquid obtained thereby, wherein the step of preparing he preparation liquid comprises (a) preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble polymer and a water-soluble salt, and (b) subjecting the first preparation liquid to a treatment of increasing a number of water-soluble salt particles, breety preparing a second preparation liquid to a first preparation liquid to a first preparation liquid.
- The process according to claim 1, wherein the water-soluble salt comprises sodium carbonate and/or sodium sulfate.
 - 3. The process according to claim 1 or 2, wherein the water-soluble polymer is one or more compounds selected

from the group consisting of acrylic acid homopolymers, acrylic acid-maleic acid copolymers and salts thereof.

 The process according to any one of claims 1 to 3, wherein the treatment of increasing a number of water-soluble salt particles comprises precipitating a water-soluble salt dissolved in the first preparation liquid.

5

20

25

- The process according to claim 4, wherein a means for precipitating the water-soluble salt dissolved in the first preparation liquid comprises adding a microcrystal-precipitating agent to the first preparation liquid.
- The process according to claim 5, wherein the microcrystal-precipitating agent is a halogenated compound of an alkali metal and/or alkaline earth metal.
 - The process according to claim 4, wherein a means for precipitating the water-soluble salt dissolved in the first preparation liquid comprises concentrating the first preparation liquid.
- 8. The process according to claim 4, wherein a means for precipitating the water-soluble salt dissolved in the first preparation liquid comprises adjusting a temperature of the first preparation liquid so that a dissolved amount of the water-soluble salt is lowered:
 - The process according to any one of claims 1 to 3, wherein the treatment of increasing a number of water-soluble salt particles comprises subjecting water-soluble salt particles in the first preparation liquid to wet grinding.
 - 10. The process sccording to any one of claims 1 to 3, wherein the treatment of increasing a number of water-soluble salt particles comprises adding to the first preparation liquid fine water-soluble salt particles which may be the same as a and/or different from the water-soluble salt in the first preparation liquid, under conditions that the fine water-soluble salt particles are capable of being present without substantially being dissolved in the first preparation liquid.
 - 11. The process according to any one of claims 1 to 3, wherein the treatment of increasing a number of water-soluble salt particles comprises two or more of the processes of claims 4 to 10.
 - 12. Particles for supporting a surfactant obtainable by spray-drying a preparation liquid comprising a water-soluble polymer and a water-soluble satt, wherein the particles for supporting a surfactant have a mode claimater of the microporous capacity distribution as determined by mercury porordineter of 1,5 μm or less, a microporous capacity of 0.3 m/Lg or more for one having a micropore diameter of from 0.01 to 3.0 μm, and a particle strength of from 15 to 100 MPa.
 - 13. The particles for supporting a surfactant according to claim 12, wherein the water-soluble polymer has a content of from 5 to 30% by weight.
 - 14. The particles for supporting a surfactant according to claim 12 or 13, further comprising a water-insoluble substance, provided that an amorphous silicate is not substantially contained.
 - 15. The particles for supporting a surfactant according to any one of claims 12 to 14, obtainable by spray-drying the preparation liquid which is obtainable by a process comprising preparing a first preparation liquid comprising a solution or slurry comprising a water-soluble polymer and a water-soluble salt, and subsequently subjecting the first preparation liquid to a treatment of increasing a number of water-soluble salt particles, thereby preparing a second preparation liquid having an increased number of water-soluble salt particles, as compared to the number of water-soluble salt particles which are present in the first preparation liquid.
- The particles for supporting a surfactant according to any one of claims 12 to 15, wherein the treatment of increasing of (1) adding a microcrystal-precipitating agent to the first preparation liquid; (2) concentrating the first preparation liquid; (3) adjusting a temperature of the first preparation liquid so that the dissolved amount of the water-soluble salt is lowered; (4) subjecting water-soluble salt particles in the first preparation liquid to wet grinding; and (5) adding to the first preparation liquid fine water-soluble salt particles which may be the same as and/or different from the water-soluble salt in the first preparation liquid on the first preparation liquid accordance in the first preparation liquid so wet grinding; and (5) adding to the first preparation liquid fine water-soluble salt particles are capable of beino gresent without substantially before dissolved in the first preparation liquid.

- 17. Particles for supporting a surfactant comprising a water-soluble polymer and a water-soluble salt, wherein at least a part of benich is a cave-in particle having a structure that there exists a hollow, namely a cave-in hole, in an inner portion thereof, and that a particle surface is opened and communicated with the hollow in the inner portion.
- 18. The particles for supporting a surfactant according to claim 17, which are obtainable by the process of any one of claims 1 to 11 or the particles of any one of claims 12 to 16.
- 19. A process for preparing delergent particles having a bulk density of from 500 to 1000 g/L, comprising the step of mixing from 10 to 100 parts by weight of a surfactant composition with 100 parts by weight of particles for supporting a surfactant obtainable by the process of any one of claims 1 to 11 or the particles of any one of claims 12 to 18.
- 20. The process according to claim 19, further comprising adding a surface coating agent.
- 21. Detergent particles having a bulk density of from 500 to 1000 g/L, wherein from 10 to 100 parts by weight of a surfactant composition is supported by 100 parts by weight of particles for supporting a surfactant obtainable by the process of any one of daims 1 to 11 or the particles of any one of claims 12 to 14.
 - 22. The detergent particles according to claim 21, wherein a surface coating agent is further added thereon.
 - 23. A detergent composition comprising the detergent particles of claim 21 or 22.

5

10

20

25

30

35

40

45

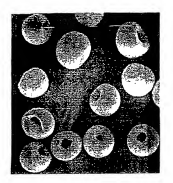


FIG. 1

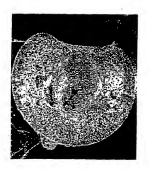


FIG. 2

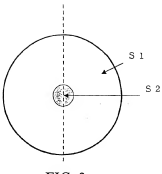


FIG. 3

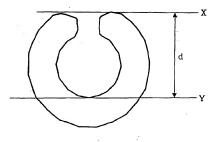


FIG. 4

INTERNATIONAL SEARCH REPORT

pcT/JP00/03856

A CLAS	SIFICATION OF SUBJECT MATTER .Cl ² CliD11/00, CliD11/02, CliC CliD3/04, CliD3/10	D17/06, C11D3/37	
According	o International Patent Classification (IPC) or to both n	ational classification and IPC	
	S SEARCHED		
Int	C11D3/04, C11D3/10	DÍ7/06, C11D3/37	
Documenta	tion searched other than minimum documentation to th	e extent that such documents are included	in the fields searched
Electronic o	tata base consulted during the international search (name	ne of data base and, where practicable, se	arch terms used)
C. DOCU	MENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.
х	EP, 221776, A2 (UNILEVER PLC), 13 May, 1987 (13.05.87),		1-11,18-23
¥	abstract; Claims; page 2, line & JP, 62-112697, A & US, 4900 & AU, 8664469, A & KR, 9004	466, A	12-16
x	EP, 289312, A2 (UNILEVER PLC), 02 November, 1988 (02.11.88),		1-11,18-23
. Y	abstract; Claims; page 3, line: & JP, 63-286496, A & US, 4820 & AU, 8815155, A & NO, 8801	441, A	12-16
x	EP, 289311, A2 (UNILEVER PLC), 02 November, 1988 (02.11.88),		1-11,18-23
Y	abstract; Claims; page 3, line: & JP, 63-286495, A & US, 4818 & AU, 8815157, A & NO, 8801	424, A	12-16
¥	US, 4547352, A (Capital City P: 15 October, 1985 (15.10.85), abstract; Claims & US, 4412978, A	roducts Company),	17-23
Furthe	r documents are listed in the continuation of Box C.	See patem family annex.	
"A" docum conside "E" carlier date "L" docum cited to special "O" docum means	categoriar of richa documents; and delinging the present that of the est which is not red to be of particular relevance considerance has published on referr the instructional filling term which reay threw doubtes on princips; claimed, see which is established by the production of the control of the con- traction in the production of the control of the con- rection is a specific of the relevancy in a mediatoristant, see, exhibition or other ent published prior to the international filling date but later protecting to an one of discinant, see, exhibition or other ent published prior to the international filling date but later protecting the control of the control of the con- parity and colors.	later document published after the inte- priority date and not conflict with it understand the principle or theory and "document of puriosity relevance," "document of puriosity relevance, the stay when the document is taken alone "or document of puriosity relevance, the considered to involve an investive site considered to involve an investive site combinated with one or more other such combinated being obvious to a person "& document member of the same passes."	te application but cited to criying the invention cannot be red to involve an inventive; claimed invention cannot be to when the document is documents, such a skilled in the art
12 5	actual completion of the international search september, 2000 (12.09.00)	Date of mailing of the international sear 26 September, 2000	
	miling address of the ISA/ nnese Patent Office	Authorized officer	

orm PCT/ISA/210 (second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

International application No. PCT/JP00/03856

(Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant pas	ssages Relevant to claim N
Y	JP, 2-255520, A (Mitajiri Kagaku Kogyo K.K.), 16 October, 1990 (16.10.90), Claims; page 3, lower right column, lines 9 to (Family: none)	17-23
Α	EP, 266863, Al (UNILEVER PLC), 11 May, 1988 (11.05.88), Claims; example 1 4 JP, 63-258998, A & AU, 8776747, A	12-16
λ	GB, 2097419, A (Colgate-Palmolive Company), 03 November, 1982 (03.11.82) 4 JP, 57-159998, A & BE, 892283, A 4 FR, 2500475, A & DE, 3206265, A1	1-23
A	JP, 4-146999, A (LION CORPORATION), 20 May; 1992 (20.05.92) (Family: none)	1-23
А	EP, 639638, Al (THE PROCTER & GAMBLE COMPAMY), 22 February, 1995 (22.02.98) & JP, 9-509436, A & WO, 95/05449, Al & CN, 1133059, A	19-23
А	EP, 421664, A2 (POHM AND HAAS COMPANY), 10 April, 1991 (10.04.91) & JP, 4-145200, A & AU, 9063264, A & CA, 2025832, A	1-23

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP00/03856

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This in	ternational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
ı. 🗆	Claims Nos.; because they relate to subject matter not required to be searched by this Authority, namely.
2.	Claims Nos.: because they relate to purce of the intermutional application that do not comply with the prescribed requirements to such an extent that no meaningful intermational search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third someroes of Rule 6.4(e).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This In	emational Searching Authority found multiple inventions in this international application, as follows:
S	ubject matters of claims 12 to 14 comprises granules for carrying a surfactant which are obtained by conventional paray drying means, and a subject matter of claim 17 comprises granules for carrying subject matter of continuous continuous continuous and continuous
ı. 🔲	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. 🛭	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. 🔲	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. 🗆	No required additional search foes were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest The additional search fees were accompanied by the applicant's protest.

Form PCT/ISA/210 (continuation of first sheet (1)) (July 1992)